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INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

1				CONTRACT (FCI)			
(51)	(51) International Patent Classification: C12N 15/86, C07K 14/16, C12N 5/10, C12N 7/04, C12N 15/49		(11) International Publication Number: (43) International Publication Date:		WO 00/15819 23 March 2000 (23.03.2000)		
(21)	International Application Number:	PCT/	US99/20675				
(22)	International Filing Date: 10 September	1999 ((10.09.1999)	Published .			
(30)	Priority Data: 60/100,022 11 September 1998 (11.0	9.199	98) US				

(60) Parent Application or Grant

60/100,063

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12 September 1998 (12.09.1998) US

(54) Title: PACKAGING CELL LINES FOR HIV-DERIVED RETROVIRAL VECTOR PARTICLES

(54) Titre: LIGNEES DE CELLULES D'ENCAPSIDATION POUR PARTICULES DE VECTEUR RETROVIRAL DERIVE DU VIH

(57) Abstract

Novel packaging cell lines useful for generating viral accessory protein independent HIV-derived retroviral vector particles, methods of constructing such packaging cell lines and methods of using the viral accessory protein independent HIV-derived retroviral vector particles are disclosed.

(57) Abrégé

L'invention concerne de nouvelles lignées de cellules d'encapsidation utiles pour produire des particules de vecteur rétroviral dérivé du VIH indépendantes de protéines accessoires virales, des procédés de mise au point de ces lignées de cellules d'encapsidation et des procédés d'utilisation des particules de vecteur rétroviral dérivé du VIH indépendantes de protéines accessoires virales.

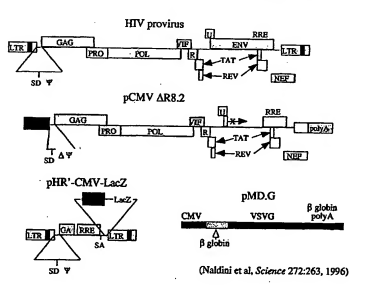
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(21) International Application Number: PCT/US9 (22) International Filing Date: 10 September 1999 (1)	•	CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT LII MC
 (30) Priority Data: 60/100,022 11 September 1998 (11.09.98 60/100,063 12 September 1998 (12.09.98 (71) Applicant: THE CHILDREN'S MEDICAL CENTER PORATION [US/US]; 300 Longwood Avenue, Bost 02115 (US). (72) Inventors: GRAY, John, T.; 48 Spring Road, West R MA 02132 (US). MULLIGAN, Richard, C.; 2 Sanc Road, Lincoln, MA 01773 (US). (74) Agents: BROOK, David, E. et al.; Hamilton, Brook, S Reynolds, P.C., Two Militia Drive, Lexington, MA (US). 	R CORton, M. Roxbury dy Pon	Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.

(54) Title: PACKAGING CELL LINES FOR HIV-DERIVED RETROVIRAL VECTOR PARTICLES



(57) Abstract

Novel packaging cell lines useful for generating viral accessory protein independent HIV-derived retroviral vector particles, methods of constructing such packaging cell lines and methods of using the viral accessory protein independent HIV-derived retroviral vector particles are disclosed.

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PACKAGING CELL LINES FOR HIV-DERIVED RETROVIRAL VECTOR PARTICLES

BACKGROUND OF THE INVENTION

Retroviral vectors based on lentiviruses, such as human immunodeficiency viruses (HIV), can infect nondividing cells, and integration of proviral DNA occurs without the need for cell division. These properties make lentiviruses attractive for gene transfer into nondividing cells, such as hepatocytes, myofibers, hematopoietic stem cells, and neurons.

However, the use of lentivirus vectors, particularly HIV vectors, particularly for gene therapy, is hampered by concern over their safety. Thus, a need for the 10 development of lentivirus vectors, particularly HIV vectors, with improved safety, particularly for gene therapy, exists.

SUMMARY OF THE INVENTION

The present invention relates to novel packaging cell lines useful for generating viral accessory protein independent lentivirus-derived, particularly HIV-derived, retroviral vector particles, to construction of such cell lines and to methods of using the accessory protein independent lentivirus-derived retroviral vector particles to introduce DNA of interest into cells (e.g, eukaryotic cells such as animal (particularly mammalian), plant or yeast cells or prokaryotic cells such as bacterial cells). In a preferred embodiment, the packaging cell lines of the present invention are stable packaging cell lines.

In one embodiment of the invention, packaging cell lines for producing a viral accessory protein independent lentivirus-derived retroviral vector particles comprise (a) a cell (e.g., mammalian cell); and (b) a retroviral nucleotide sequence in the cell which comprises a coding sequence for lentivirus gagpol, wherein said coding sequence has

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been codon optimized by mutagenisis to improve expression of the lentivirus gagpol proteins.

In second embodiment of the invention, packaging cell lines for producing a viral accessory protein independent lentivirus-derived retroviral vector particles comprise (a) a cell (e.g., mammalian cell); (b) a first retroviral nucleotide sequence in the cell which comprises a coding sequence for lentivirus gagpol, wherein said coding sequence has been codon optimized by mutagenisis to improve expression of the lentivirus gagpol proteins; and (c) a second retroviral nucleotide sequence in the cell which comprises the coding sequence for a heterologous envelope protein.

In a third embodiment of the invention, packaging cell lines for producing a viral accessory protein independent lentivirus-derived retroviral vector particles comprise (a) a cell (e.g., mammalian cell); (b) a first retroviral nucleotide sequence in the cell which comprises a coding sequence for lentivirus gagpol, wherein said coding sequence has been codon optimized by mutagenisis to improve expression of the lentivirus gagpol proteins; (c) a second retroviral nucleotide sequence in the cell which comprises the coding sequence for a heterologous envelope protein; and (d) a third retroviral nucleotide sequence which comprises a DNA sequence of interest and lentivirus cisacting sequences required for packaging, reverse transcription and integration.

In a fourth embodiment of the invention, packaging cell lines for producing a viral accessory protein independent lentivirus-derived retroviral vector particles comprise (a) a cell (e.g., mammalian cell); (b) a retroviral nucleotide sequence in the cell which comprises a coding sequence for lentivirus gagpol, wherein said coding sequence has been codon optimized by mutagenisis to improve expression of the lentivirus gagpol proteins; and (c) a retroviral nucleotide sequence which comprises a DNA sequence of interest and lentivirus cis-acting sequences required for packaging, reverse transcription and integration.

In a fifth embodiment of the invention, packaging cell lines for producing a viral accessory protein independent HIV-derived retroviral vector particles comprise (a) a cell

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(e.g., mammalian cell); and (b) a retroviral nucleotide sequence in the cell which comprises a coding sequence for HIV gagpol, wherein said coding sequence has been codon optimized by mutagenisis to improve expression of the HIV gagpol proteins.

In sixth embodiment of the invention, packaging cell lines for producing a viral accessory protein independent HIV-derived retroviral vector particles comprise (a) a cell (e.g., mammalian cell); (b) a first retroviral nucleotide sequence in the cell which comprises a coding sequence for HIV gagpol, wherein said coding sequence has been codon optimized by mutagenisis to improve expression of the HIV gagpol proteins; and (c) a second retroviral nucleotide sequence in the cell which comprises the coding sequence for a heterologous envelope protein.

In a seventh embodiment of the invention, packaging cell lines for producing a viral accessory protein independent HIV-derived retroviral vector particles comprise (a) a cell (e.g., mammalian cell); (b) a first retroviral nucleotide sequence in the cell which comprises a coding sequence for HIV gagpol, wherein said coding sequence has been codon optimized by mutagenisis to improve expression of the HIV gagpol proteins; (c) a second retroviral nucleotide sequence in the cell which comprises the coding sequence for a heterologous envelope protein; and (d) a third retroviral nucleotide sequence which comprises a DNA sequence of interest and HIV cis-acting sequences required for packaging, reverse transcription and integration.

In a eighth embodiment of the invention, packaging cell lines for producing a viral accessory protein independent HIV-derived retroviral vector particles comprise (a) a cell (e.g., mammalian cell); (b) a retroviral nucleotide sequence in the cell which comprises a coding sequence for HIV gagpol, wherein said coding sequence has been codon optimized by mutagenisis to improve expression of the HIV gagpol proteins; and (c) a retroviral nucleotide sequence which comprises a DNA sequence of interest and HIV cis-acting sequences required for packaging, reverse transcription and integration.

Alternatively, each of the packaging cell lines described herein can be produced using (1) a retroviral nucleotide sequence which comprises a codon optimized gag

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coding sequence and (2) a retroviral nucleotide sequence which comprises a codon optimized pol coding sequence, in place of the retroviral nucleotide sequence which comprises a codon optimized gagpol coding sequence.

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In a particular embodiment, the heterologous envelope protein is the G glycoprotein of vesicular stomatitis virus (VSV G). In another embodiment, the heterologous envelope protein is the amphotropic envelope of the Moloncy leukemia virus (MLV).

Cell lines for producing a viral accessory protein independent lentivirus-derived retroviral vector particles are produced by transfecting host cells (e.g., mammalian host cells) with a plasmid comprising a DNA sequence which encodes lentivirus gagpol proteins, wherein said DNA sequence has been codon optimized by mutagenisis to improve expression of the lentivirus gagpol proteins. Depending upon the particular cell line being produced, the host cells are also co-transfected with a plasmid comprising a DNA sequence which encodes a heterologous envelope protein, or a plasmid comprising a DNA sequence of interest and lentivirus cis-acting sequences required for packaging, reverse transcription and integration, or both of these plasmids. Alternatively, host cells are transfected with a plasmid comprising a codon optimized DNA sequence encoding a lentivirus gag protein and a plasmid comprising a codon optimized DNA sequence encoding a lentivirus pol protein, in place of the plasmid comprising a codon optimized DNA sequence encoding both lentivirus gagpol proteins.

Cell lines for producing a viral accessory protein independent HIV-derived retroviral vector particles are produced by co-transfecting host cells (e.g., mammalian host cells) with a plasmid comprising a DNA sequence which encodes HIV gagpol proteins, wherein said DNA sequence has been codon optimized by mutagenisis to improve expression of the HIV gagpol proteins. Depending upon the particular cell line being produced, the host cells are also co-transfected with a plasmid comprising a DNA sequence which encodes a heterologous envelope protein, or a plasmid comprising a DNA sequence of interest and HIV cis-acting sequences required for packaging, reverse

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transcription and integration, or both of these plasmids. Alternatively, host cells are transfected with a plasmid comprising a codon optimized DNA sequence encoding a HIV gag protein and a plasmid comprising a codon optimized DNA sequence encoding a HIV pol protein, in place of the plasmid comprising a codon optimized DNA sequence encoding both HIV gagpol proteins.

The present invention also relates to methods of producing viral accessory protein independent lentivirus-derived retroviral vector particles, comprising cotransfecting host cells (e.g., mammalian host cells) with (a) a first plasmid comprising a DNA sequence which encodes lentivirus gagpol proteins, wherein said DNA sequence has been codon optimized by mutagenisis to improve expression of the lentivirus gagpol proteins; (b) a second plasmid comprising a DNA sequence which encodes a heterologous envelope protein; and (c) a third plasmid comprising a DNA sequence of interest and lentivirus cis-acting sequences required for packaging, reverse transcription and integration. Alternatively, host cells are transfected with a plasmid comprising a codon optimized DNA sequence encoding a lentivirus gag protein and a plasmid comprising a codon optimized DNA sequence encoding a lentivirus pol protein, in place of the first plasmid comprising a codon optimized DNA sequence encoding both lentivirus gagpol proteins.

In a particular embodiment, the invention relates to methods of producing viral accessory protein independent HIV-derived retroviral vector particles, comprising cotransfecting host cells (e.g., mammalian host cells) with (a) a first plasmid comprising a DNA sequence which encodes HIV gagpol proteins, wherein said DNA sequence has been codon optimized by mutagenisis to improve expression of the HIV gagpol proteins; (b) a second plasmid comprising a DNA sequence which encodes a heterologous envelope protein; and (c) a third plasmid comprising a DNA sequence of interest and HIV cis-acting sequences required for packaging, reverse transcription and integration. Alternatively, bost cells are transfected with a plasmid comprising a codon optimized DNA sequence encoding a HIV gag protein and a plasmid comprising a

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codon optimized DNA sequence encoding a HIV pol protein, in place of the first

plasmid comprising a codon optimized DNA sequence encoding both HIV gagpol

proteins.

The present invention also relates to viral accessory protein-independent retroviral particles produced by or obtainable by (obtained by) the methods described herein.

The present invention further relates to isolated DNA encoding a codon optimized lentivirus gagpol, isolated DNA encoding the gag coding region of a codon optimized lentivirus gagpol, and isolated DNA encoding the pol coding region of a codon optimized lentivirus gagpol. In a particular embodiment, the present invention relates to isolated DNA encoding a codon optimized HIV gagpol, isolated DNA encoding the gag coding region of a codon optimized HIV gagpol, and isolated DNA encoding the pol coding region of a codon optimized HIV gagpol.

The packaging cell lines and viral particles of the present invention can be used

for gene therapy or gene replacement with improved safety. The packaging cell lines
and viral particles of the present invention can also be used in development and
production of vaccines, and in production of biochemical reagents. Gene therapy
vectors produced with the cell lines of the present invention are expected to be valuable
medical therapeutics.

20 BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 is a schematic diagram of an expression cassette containing the codon optimized gagpol genes. The DNA was constructed in multiple segments, which are indicated at the top as 1/3, 2/3, 3/3 (A, B, C and D) and HIN. Restriction sites used to assemble the cloned segments are indicated above the kilobasepair (Kb) ruler. Below the ruler are multiple features showing the location of the human cytomegalovirus (CMV) promoter, human betaglobin sequences (Bglobin), mRNA sequences (thinner line represents intronic sequence), the gag and pol open reading frames, the individual

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proteolytic fragment coding sequences (p17_MA, p24_CA, p7, p6, PR, p51_RT, RNaseH and integrase (IN)) and each synthetic oligonucleotide used in the assembly process (multiple adjacent open arrows).

Figure 2 is a table which depicts codon usage frequencies in genes which are highly expressed and in the codon optimized gagpol open reading frame of the HIV packaging construct described herein.

Figure 3 is a schematic representation of the HIV provirus and a three-plasmid expression system used for generating a pseudotyped HIV-based vector by transient transfection as described in Naldini *et al.*, *Science*, *272*:263-267 (1996).

Figure 4 is a list of some characteristics relating to the HIV Rev protein.

Figure 5 is a list of some points relating to codon optimization of HIV gagpol.

Figure 6 is a partial DNA sequence of HIV gag (SEQ ID NO: 1), showing inactivation of inhibitory sequences as described in Schwartz, S. et al., J. Virol.,

66(12):7176-7182 (1992).

Figure 7 a plot of the %(G+C) content of wildtype IIIV gagpol sequences and theoretically codon optimized HIV gagpol sequences. The percent of bases, either G or C, was calculated for a 30 nucleotide moving window for the entire length of the gagpol gene, and the value plotted versus nucleotide position. Diamonds = HIV gagpol sequences; squares = full optimal back-translation for gag open reading frame; triangles = full optimal back-translation for pol open reading frame; CO = codon optimized.

Figures 8A-8E depict the alignment of the nucleotide sequences and predicted amino acid sequences for the gag coding region of a wildtype HIV gagpol and a codon optimized HIV gagpol. "NL4-3 genbank SEQ" indicates the nucleotide sequence (SEQ ID NO:2) and predicted amino acid sequence (SEQ ID NO:3) for the gag coding region of a wildtype HIV gagpol. "pHDMHgpm2.seq" indicates the nucleotide sequence (SEQ ID NO:4) and predicted amino acid sequence (SEQ ID NO:5) for the gag coding region

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of a codon optimized HIV gagpol. The "NL4-3 genbank.SEQ" sequences are publicly available at the NIH GenBank sequence repository (Accesssion No. M19921).

Figures 9A-9L depict the alignment of the nucleotide sequences and predicted amino acid sequences for the *pol* coding region of a wildtype HIV *gagpol* and a codon optimized HIV *gagpol*. "NL4-3 genbank.SEQ" indicates a nucleotide sequence (SEQ ID NO:6) and a predicted amino acid sequence (SEQ ID NO:7) for the *pol* coding region of a wildtype HIV *gagpol* available in the NIH GenBank sequence repository (Accesssion No. M19921). The nucleotide and amino acid sequences for the *pol* coding region available in the GenBank sequence repository contain two sequence errors, which are indicated in Figures 9A-9L with shading. "pNL4-3.seq" indicates the correct nucleotide sequence (SEQ ID NO:8) and predicted amino acid sequence (SEQ ID NO:9) for the *pol* coding region of a wildtype HIV *gagpol*. "pHDMHgpm2.seq" indicates the nucleotide sequence (SEQ ID NO:10) and predicted amino acid sequence (SEQ ID NO:11) for the *pol* coding region of a codon optimized HIV *gagpol*.

Figures 10A-10D depict the DNA sequence (SEQ ID NO:12) for pHDMHgpm2. The CMV enhancer/promoter is at nucleotides 97 to 679, human betaglobin sequences (Bglobin) are at nucleotides 761 to 864, 865 to 1303 and 5710 to 6469 (end of Bglobin is at nucleotides 6445 to 6469), mRNA sequences are at nucleotides 680 to 778 and 1255 to 5921, SV40 origin of replication is at nucleotides 8796 to 8908, beta-lactamase (bla) coding region is at nucleotides 6709 to 7569, intron sequences are at nucleotides 779 to 1254, the codon optimized *gag* coding region is at nucleotides 1318 to 2820, the codon optimized *pol* coding region is at nucleotides 2619 to 5624 and the poly A site is at nucleotides 5897 to 5921.

Figure 11 is a circular map of plasmid pHDMHgpm2.

25 DETAILED DESCRIPTION OF THE INVENTION

The present invention relates to novel packaging cell lines useful for generating viral accessory protein independent lentivirus-derived, particularly HIV-derived,

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retroviral vector particles, to construction of such cell lines and to methods of using the accessory protein independent lentivirus-derived retroviral vector particles to introduce DNA of interest into cells (e.g, eukaryotic cells such as animal (particularly mammalian), plant or yeast cells or prokaryotic cells such as bacterial cells). In a particular embodiment, the packaging cell lines of the present invention are stable packaging cell lines.

The cell lines are engineered to express the lentivirus proteins necessary for virus particle formation (gagpol proteins), without containing DNA sequences from lentivirus accessory proteins (tat, vif, vpr, vpu, nef and rev proteins and Rev response element (RRE)). Additionally, no viral sequences (such as cis-acting elements termed constitutive transport elements (CTEs)) will be expressed as RNA of any kind. DNA sequences for lentivirus gagpol are codon optimized by extensively mutagenizing the sequences to improve expression and to reduce the risk of recombination between transfer vector sequences and gagpol messenger RNA. This greatly improves the safety of virus preparations generated from these cell lines. In a particular embodiment, the DNA sequences for lentivirus gagpol are not codon optimized in the overlap region between the gag and pol sequences and in cis-acting signals necessary for translation of

Examples of lentiviruses include human immunodeficiency viruses (e.g., HIV-1, HIV-2, HIV-3), bovine lentiviruses (e.g., bovine immunodeficiency viruses, bovine immunodeficiency-like viruses, Jembrana disease viruses), equine lentiviruses (e.g., equine infectious anemia viruses), feline lentiviruses (e.g., feline immunodeficiency viruses, panther lentiviruses, puma lentiviruses), ovine/caprine lentiviruses (e.g., Brazilian caprine lentiviruses, caprine arthritis-encephalitis viruses, Maedi-Visna viruses, Maedi-Visna-like viruses, Maedi-Visna-related viruses, ovine lentiviruses, Visna lentiviruses), Simian AIDS retroviruses (e.g., human T-cell lymphotropic virus type 4), simian immunodeficiency viruses, simian-human immunodeficiency viruses, human lymphotrophic viruses (e.g., type III), simian T-cell lymphotrophic viruses.

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In another embodiment, cell lines are engineered to express the HIV proteins necessary for virus particle formation (gagpol proteins), without containing DNA sequences from HIV accessory proteins (tat, vif, vpr, vpu, nef and rev proteins and Rev response element (RRE)). Additionally, no viral sequences (such as cis-acting elements termed constitutive transport elements (CTEs)) will be expressed as RNA of any kind. DNA sequences for a HIV gagpol are codon optimized by mutagenesis to improve expression and to reduce the risk of recombination between transfer vector sequences and gagpol messenger RNA. In a particular embodiment, the DNA sequences for HIV gagpol are not codon optimized in the overlap region between the gag and pol sequences and in cis-acting signals necessary for translation of pol.

Alternatively, each of the packaging cell lines described herein can be produced using (1) a nucleotide sequence which comprises a codon optimized gag coding sequence and (2) a nucleotide sequence which comprises a codon optimized pol coding sequence, in place of the nucleotide sequence which comprises a codon optimized gagpol coding sequence. In this embodiment, the gag and pol coding sequences can be completely codon optimized

Benefits of the present invention include the removal of potentially harmful lentivirus accessory proteins and other viral sequences, and the reduction of the risk of recombination to produce replication competent virus.

Packaging cell lines for producing a viral accessory protein independent lentivirus-derived retroviral vector particles comprise a mammalian cell and a retroviral nucleotide sequence comprising a coding sequence for a lentivirus gagpol which has been codon optimized. In a particular embodiment the packaging cell lines further comprise a retroviral nucleotide sequence comprising a coding sequence for a heterologous envelope protein. In a second embodiment, the packaging cell lines further comprise a retroviral nucleotide sequence comprising a coding sequence for a heterologous envelope protein and a retroviral nucleotide sequence which comprises a DNA sequence of interest and HIV cis-acting sequences required for packaging, reverse

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transcription and integration. In third embodiment, the packaging cell lines further comprise a retroviral nucleotide sequence which comprises a DNA sequence of interest and HIV cis-acting sequences required for packaging, reverse transcription and integration. Alternatively, the packaging cell lines of the present invention comprise a retroviral nucleotide sequence which comprises a codon optimized gag coding sequence and (2) a retroviral nucleotide sequence which comprises a codon optimized pol coding sequence, in place of the retroviral nucleotide sequence which comprises a codon optimized gagpol coding sequence.

The coding sequence(s) for lentivirus gagpol which has (have) been codon optimized results in improved expression of the lentivirus gagpol proteins and reduces the risk of recombination between the transfer vector and gagpol messenger RNA. Codon optimization of the coding sequence(s) for lentivirus gagpol was obtained by mutagenizing for each particular amino acid residue, specific nucleic acid bases in a codon for the particular amino acid residue to a nucleic acid base which is present in a codon which occurs at a high frequency in genes which are highly expressed for the same amino acid residue. In a particular embodiment, the resulting optimized codon also does not cause introduction of mRNA splicing signals into the codon optimized sequence. Thus, in a particular embodiment, codon optimization of the coding sequence(s) for lentivirus gagpol is obtained by mutagenizing for each particular amino acid residue, specific nucleic acid bases in a codon for the particular amino acid residue to a nucleic acid base that is present in a codon which (1) occurs at a high frequency in genes which are highly expressed for the same amino acid residue and (2) does not cause introduction of mRNA splicing signals into the codon optimized sequence. Codon optimization typically results in the removal of nucleic acid base A-rich instability elements.

In a particular embodiment, the coding sequence for a HIV gagpol (pNL4-3; available through the AIDS repository, NIH; Adachi et al., J. Virol., 59:284-291 (1986)) has been codon optimized to improve translational efficiency of the HIV gagpol

proteins and reduce the risk of recombination between the transfer vector and HIV gagpol messenger RNA. Two hundred thirty-seven base pairs (237 bp) consisting of the gag pol overlap and cis-acting signals necessary for translation of pol (nucleotides 2583 to 2819 of SEQ ID NO: 12) were not optimized. The HIV gagpol sequence obtained—using the codon optimization process does not differ at the amino acid level from the wildtype HIV gagpol sequence, but differs at the nucleotide level from the HIV gagpol sequence. A codon optimized HIV gag sequence is shown in Figures 8A-8E (pHDMHgpm2.seq) (SEQ ID NO:4). A codon optimized HIV pol sequence is shown in Figures 9A-9L (pHDMHgpm2.seq) (SEQ ID NO:10).

A plasmid comprising DNA sequences which encode codon optimized lentivirus gagpol proteins is also referred to herein as a packaging construct. This plasmid includes a promoter which drives the expression of the gagpol proteins, such as the human cytomegalovirus (hCMV) immediate early promoter. This plasmid is defective for the production of the viral envelope and accessory proteins tat, vif, vpr, vpu, nef and rev and the Rev response element (RRE). The packaging construct also does not contain viral sequences which are transcribed into mRNA, such as constitutive transport elements (CTEs).

A packaging construct comprising a codon optimized HIV gagpol is depicted in Figure 1 and in Figure 11. Figures 10A-10D depict the DNA sequence (SEQ ID NO:12) for the packaging construct pHDMHgpm2. This packaging construct (pHDMHgpm2) was constructed as follows: Plasmid pMDA.HIVgp mam was generated by chemical synthesis and PCR assembly (which is described in, for example, Stemmer et al., Gene, 164:49-53 (1995)) of 215 different oligonucleotides. The DNA sequence for pMDA.HIVgp mam is the same as the DNA sequence for pMDA.HIVgp jtg except for 4.3 kb which was codon optimized using the DNAStar program (LaserGene, Madison, WI). Two hundred thirty-seven base pairs (237 bp) consisting of the gag pol overlap and cis-acting signals necessary for translation of pol (nucleotides 2583 to 2819 of SEQ ID NO: 12) were not optimized due to dual reading

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frame constraints. A Nsil site 5' of IN was preserved to aid fusion with wildtype sequences. Several single or double base pair silent mutations were introduced either to prevent potential splice donors and acceptors, or by the synthesis process. pMDA.HIVgp jtg was derived from HIV-1 strain NL4-3. The protease mutation that is present in the NL4-3 NIH GenBank sequence was then repaired (Figure 9B), changing the nucleotide present at position 2948 of SEQ ID NO:12 from a "G" to a "C", thereby producing the codon present at nucleotide positions 2948 to 2950 of SEQ ID NO:12 which encodes an arginine instead of the glycine present in the NL4-3 GenBank amino acid sequence. The resulting plasmid was named pMDHgpmam. The EcoRI-HindIII fragment of pMDHgpmam was inserted into pHDM2b, a high copy version of the pMD vector (Ory, D. et al., Proc. Natl. Acad. Sci. USA, 93(21):11400-11406 (1996)), to produce plasmid pHDMHgpm. The sequencing mutation that is present in the RNase domain of the NL4-3 NIH GenBank sequence was repaired (Figure 9H), changing the codon present at nucleotide positions 4724 to 4726 of SEQ ID NO:12 from "GGG" to "AAG", thereby producing a codon encoding a lysine instead of the glycine present in the NL4-3 GenBank amino acid sequence. The resulting plasmid was named pHDMHgpm2. Codon usage frequencies in the codon optimized gagpol open reading frame of the packaging construct pHDMHgpm2 are shown in Figure 2.

As used herein, a heterologous envelope protein permits pseudotyping of particles generated by the packaging construct and includes the G glycoprotein of vesicular stomatitis virus (VSV G) and the amphotropic envelope of the Moloney leukemia virus (MLV). A plasmid comprising a DNA sequence which encodes a heterologous envelope protein is also referred to herein as an envelope coding plasmid.

The terms "mammal" and "mammalian", as used herein, refer to any vertebrate animal, including monotremes, marsupials and placental, that suckle their young and either give birth to living young (eutharian or placental mammals) or are egg-laying (metatharian or nonplacental mammals). Examples of mammalian species include

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humans and other primates (e.g., monkeys, chimpanzees), rodents (e.g., rats, mice, guinea pigs) and ruminents (e.g., cows, pigs, horses).

Examples of mammalian cells include human (such as HeLa cells, 293T cells, NIH 3T3 cells), bovine, ovine, porcine, murine (such as embryonic stem cells), rabbit and monkey (such as COS1 cells) cells. The cell may be a non-dividing cell (including hepatocytes, myofibers, hematopoietic stem cells, neurons) or a dividing cell. The cell may be an embryonic cell, bone marrow stem cell or other progenitor cell. Where the cell is a somatic cell, the cell can be, for example, an epithelial cell, fibroblast, smooth muscle cell, blood cell (including a hematopoietic cell, red blood cell, T-cell, B-cell, etc.), tumor cell, cardiac muscle cell, macrophage, dendritic cell, neuronal cell (e.g., a glial cell or astrocyte), or pathogen-infected cell (e.g., those infected by bacteria, viruses, virusoids, parasites, or prions).

Typically, cells isolated from a specific tissue (such as epithelium, fibroblast or hematopoietic cells) are categorized as a "cell-type." The cells can be obtained commercially or from a depository or obtained directly from an animal, such as by biopsy. Alternatively, the cell need not be isolated at all from the animal where, for example, it is desirable to deliver the virus to the animal in gene therapy.

To produce the cell lines of the present invention for producing a viral accessory protein independent lentivirus-derived retroviral vector particles, mammalian host cells are co-transfected with (a) a first plasmid comprising DNA sequence which encode lentivirus gagpol proteins, wherein said DNA sequence has been codon optimized by mutagenisis, as described above, to improve expression of the lentivirus gagpol proteins; and (2) a second plasmid comprising a DNA sequence which encodes a heterologous envelope protein, or a retroviral nucleotide sequence which comprises a DNA sequence of interest and lentivirus cis-acting sequences required for packaging, reverse transcription and integration, or both, under conditions appropriate for transfection of the cells.

-15-

In a particular embodiment, to produce the cell lines of the present invention for producing viral accessory protein independent HIV-derived retroviral vector particles mammalian host cells were cotransfected with (a) a first plasmid comprising DNA sequence which encode HIV gagpol proteins, wherein said DNA sequence has been codon optimized by mutagenisis, as described above, to improve expression of the HIV gagpol proteins; and (2) a second plasmid comprising a DNA sequence which encodes a heterologous envelope protein, or a retroviral nucleotide sequence which comprises a DNA sequence of interest and HIV cis-acting sequences required for packaging, reverse transcription and integration, or both, under conditions appropriate for transfection of the cells.

Virus stocks consisting of viral accessory protein independent lentivirus-derived, particularly HIV-derived, retroviral vector particles of the present invention are produced by maintaining the transfected cells under conditions suitable for virus production (e.g., in an appropriate growth media and for an appropriate period of time). Such conditions, which are not critical to the invention, are generally known in the art. See, e.g., Sambrook et al., Molecular Cloning: A Laboratory Manual, Second Edition, Cold Spring Harbor University Press, New York (1989); Ausubel et al., Current Protocols in Molecular Biology, John Wiley & Sons, New York (1998); U.S. Patent No. 5,449,614; and U.S. Patent No. 5,460,959, the teachings of which are incorporated herein by reference.

To generate viral accessory protein independent lentivirus-derived retroviral vector particles, mammalian host cells can be co-transfected with (a) a first plasmid comprising DNA sequence which encode lentivirus gagpol proteins, wherein said DNA sequence has been codon optimized by mutagenisis, as described above, to improve expression of the lentivirus gagpol proteins; (b) a second plasmid comprising a DNA sequence which encodes a heterologous envelope protein; and (c) a third plasmid comprising a DNA sequence of interest and lentivirus cis-acting sequences required for packaging, reverse transcription and integration. Alternatively, mammalian cells are

transfected with a plasmid comprising a codon optimized DNA sequence encoding a lentivirus gag protein and a plasmid comprising a codon optimized DNA sequence encoding a lentivirus pol protein, in place of the first plasmid comprising a codon optimized DNA sequence encoding both lentivirus gagpol proteins. Alternatively, — mammalian host cells are transfected with a plasmid comprising a codon optimized DNA sequence encoding a lentivirus gag protein and a plasmid comprising a codon optimized DNA sequence encoding a lentivirus pol protein, in place of the first plasmid comprising a codon optimized DNA sequence encoding both lentivirus gagpol proteins.

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In a particular embodiment, the invention relates to methods of producing viral accessory protein independent HIV-derived retroviral vector particles, comprising cotransfecting mammalian host cells with (a) a first plasmid comprising DNA sequence which encode HIV gagpol proteins, wherein said DNA sequence has been codon optimized by mutagenisis, as described above, to improve expression of the HIV gagpol proteins; (b) a second plasmid containing a DNA sequence which encodes a heterologous envelope protein; and (c) a third plasmid comprising a DNA sequence of interest and HIV cis-acting sequences required for packaging, reverse transcription and integration. Alternatively, mammalian host cells are transfected with a plasmid comprising a codon optimized DNA sequence encoding a HIV gag protein and a plasmid comprising a codon optimized DNA sequence encoding a HIV pol protein, in place of the first plasmid comprising a codon optimized DNA sequence encoding both HIV gagpol proteins.

Virus particles produced by the methods described herein, using a codon optimized HIV packaging construct produced as described herein, were compared by Western analysis with virus particles produced as described in Naldini et al., Science, 272:263-267 (1996), using the packaging construct plasmid pCMVAR8.2. Both the immunological reactivity and the proteolytic processing were confirmed to be indistinguishable.

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A plasmid comprising a DNA sequence of interest and HIV cis-acting sequences required for packaging, reverse transcription and integration is also referred to herein as a transfer vector. A transfer vector, as used herein, refers to a vehicle which is used to introduce a DNA of interest into a eurkaryotic cell, particularly a mammalian cell.

Figure 3 depicts an example of a transfer vector.

DNA sequence of interest, as used herein, include all or a portion of a gene or genes encoding a nucleic acid product whose expression in a cell or a mammal is desired. In a particular embodiment, the nucleic acid product is a heterologous therapeutic protein. Examples of therapeutic proteins include antigens or immunogens, such as a polyvalent vaccine, cytokines, tumor necrosis factor, interferons, interleukins, adenosine dearninase, insulin, T-cell receptors, soluble CD4, growth factors, such as epidermal growth factor, human growth factor, insulin-like growth factors, fibroblast growth factors), blood factors, such as Factor VIII, Factor IX, cytochrome b, glucocerebrosidase, ApoE, ApoC, ApoAl, the LDL receptor, negative selection markers or "suicide proteins", such as thymidine kinase (including the HSV, CMV, VZV TK), anti-angiogenic factors, Fc receptors, plasminogen activators, such as t-PA, u-PA and streptokinase, dopamine, MHC, tumor suppressor genes such as p53 and Rb, monoclonal antibodies or antigen binding fragments thereof, drug resistance genes, ion channels, such as a calcium channel or a potassium channel, adrenergic receptors, hormones (including growth hormones) and anti-cancer agents. In another embodiment, the nucleic acid product is a gene product to be expressed in a cell or a mammal and which product is otherwise defective or absent in the cell or mammal. For example, the nucleic acid product can be a functional gene(s) which is defective or absent in the cell or mammal.

DNA sequence of interest includes DNA sequences (control sequences) which are necessary to drive the expression of the gene or genes. The control sequences are operably linked to the gene. The term "operably linked", as used herein, is defined to mean that the gene is linked to control sequences in a manner which allows expression

-18-

of the gene (or the nucleic acid sequence). Generally, operably linked means contiguous.

Control sequences include a transcriptional promoter, an optional operator sequence to control transcription, a sequence encoding suitable mRNA ribosomal — binding sites and sequences which control termination of transcription and translation. In a particular embodiment, a recombinant gene encoding a desired nucleic acid product can be placed under the regulatory control of a promoter which can be induced or repressed, thereby offering a greater degree of control with respect to the level of the product produced.

As used herein, the term "promoter" refers to a sequence of DNA, usually upstream (5') of the coding region of a structural gene, which controls the expression of the coding region by providing recognition and binding sites for RNA polymerase and other factors which may be required for initiation of transcription. Suitable promoters are well known in the art. Exemplary promoters include the SV40, CMV and human elongation factor (EFI) promoters. Other suitable promoters are readily available in the art (see, e.g., Ausubel et al., Current Protocols in Molecular Biology, John Wiley & Sons, Inc., New York (1998); Sambrook et al., Molecular Cloning: A Laboratory Manual, 2nd edition, Cold Spring Harbor University Press, New York (1989); and U.S. Patent No. 5,681,735).

A DNA sequence of interest can be isolated from nature, modified from native sequences or manufactured de novo, as described in, for example, Ausubel et al., Current Protocols in Molecular Biology, John Wiley & Sons, New York (1998); and Sambrook et al., Molecular Cloning: A Laboratory Manual, 2nd edition, Cold Spring Harbor University Press, New York. (1989). DNA sequences can be isolated and fused together by methods known in the art, such as exploiting and manufacturing compatible cloning or restriction sites.

The packaging cell lines and viral particles of the present invention can be used, in vitro, in vivo and ex vivo, to introduce DNA of interest into a eukaryotic cell (e.g., a

mammalian cell) or a mammal (e.g., a human or other mammal or vertebrate). The cells can be obtained commercially or from a depository or obtained directly from a mammal, such as by biopsy. The cells can be obtained from a mammal to whom they will be returned or from another/different mammal of the same or different species. For example, using the packaging cell lines or viral particles of the present invention, DNA of interest can be introduced into nonhuman cells, such as pig cells, which are then introduced into a human. Alternatively, the cell need not be isolated from the mammal where, for example, it is desirable to deliver vial particles of the present invention to the mammal in gene therapy.

Ex vivo therapy has been described, for example, in Kasid et al., Proc. Natl. Acad. Sci. USA, 87:473 (1990); Rosenberg et al., N. Engl. J. Med., 323:570 (1990); Williams et al., Nature, 310:476 (1984); Dick et al., Cell, 42:71 (1985); Keller et al., Nature, 318:149 (1985); and Anderson et al., United States Patent No. 5,399,346.

Methods for administering (introducing) viral particles directly to a mammal are generally known to those practiced in the art. For example, modes of administration include parenteral, injection, mucosal, systemic, implant, intraperitoneal, oral, intradermal, transdermal (e.g., in slow release polymers), intramuscular, intravenous including infusion and/or bolus injection, subcutaneous, topical, epidural, etc. Viral particles of the present invention can, preferably, be administered in a pharmaceutically acceptable carrier, such as saline, sterile water, Ringer's solution, and isotonic sodium chloride solution.

The dosage of a viral particle of the present invention administered to a mammal, including frequency of administration, will vary depending upon a variety of factors, including mode and route of administration; size, age, sex, health, body weight and diet of the recipient mammal; nature and extent of symptoms of the disease or disorder being treated; kind of concurrent treatment, frequency of treatment, and the effect desired.

-20-

The teachings of all the articles, patents, patent applications and GenBank sequences cited herein are incorporated by reference in their entirety.

While this invention has been particularly shown and described with references to preferred embodiments thereof, it will be understood by those skilled in the art that various changes in form and details may be made therein without departing from the spirit and scope of the invention as defined by the appended claims.

Claims

-21-

CLAIMS

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 A packaging cell line for producing a viral accessory protein independent HIVderived retroviral vector particle comprising:

5 a) a mammalian cell;

What is claimed is:

- b) a first retroviral nucleotide sequence in the cell which comprises a coding sequence for a HIV gagpol, wherein said coding sequence has been mutagenized to improve expression of the HIV gagpol proteins:
- a second retroviral nucleotide sequence in the cell which comprises the coding sequence for a heterologous envelope protein; and
- a third retroviral nucleotide sequence in the cell which comprises a DNA sequence of interest and HIV cis-acting sequences required for packaging, reverse transcription and integration.

A packaging cell line of Claim 1 wherein the heterologous envelope protein is
 the G glycoprotein of vesicular stomatitis virus (VSV G).

- A packaging cell line of Claim 1 wherein the heterologous envelope protein is the amphotropic envelope of the Moloney leukemia virus.
- A packaging cell line of Claim I wherein the DNA sequence of interest encodes
 a heterologous therapeutic protein.
- 20 5. A packaging cell line comprising:
 - a) a mammalian cell;

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	. 5		 a first retroviral nucleotide sequence in the cell which comprises a coding sequence for a HIV gagpol, wherein said coding sequence has been mutagenized to improve expression of the HIV gagpol proteins; and a second retroviral nucleotide sequence in the cell which comprises a DNA sequence of interest and HIV cis-acting sequences required for
15			packaging, reverse transcription and integration.
20		6.	A packaging cell line of Claim 5 wherein the DNA sequence of interest encodes a heterologous therapeutic protein.
25	10	7.	A packaging cell line comprising: a) a mammalian cell; b) a first retroviral nucleotide sequence in the cell which comprises a coding sequence for a HIV gagpol, wherein said coding sequence has
30	15		been mutagenized to improve expression of the HIV gagpol proteins; and c) a second retroviral nucleotide sequence in the cell which comprises the coding sequence for a heterologous envelope protein.
35		8.	A method of producing a packaging cell line for producing a viral accessory protein independent HIV-derived retroviral vector particle, comprising cotransfecting mammalian host cells with: a) a first plasmid comprising a DNA sequence which encodes HIV gagpol
40	20		a) a first plasmid comprising a DNA sequence which encodes in v gagpor proteins, wherein said DNA sequence has been mutagenized to improve expression of the HIV gag and pol proteins; b) a second plasmid comprising a DNA sequence which encodes a
45			heterologous envelope protein; and

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10	· :	 a third plasmid comprising a DNA sequence of interest and HIV cis- acting sequences required for packaging, reverse transcription and integration.
15	9. 5	A method of Claim 8 wherein the heterologous envelope protein is the G glycoprotein of vesicular stomatitis virus (VSV G).
20	10.	A method of Claim 8 wherein the heterologous envelope protein is the amphotropic envelope of the Moloney leukemia virus.
25	11.	A method of Claim 8 wherein the DNA sequence of interest is a heterologous therapeutic protein.
	10 12.	A method of producing a viral accessory protein independent HIV-derived retroviral vector particle comprising co-transfecting mammalian host cells with:
30		 a first plasmid comprising a DNA sequence which encodes HIV gagpol proteins, wherein said DNA sequence has been mutagenized to improve expression of the HIV gagpol proteins;
35	15	b) a second plasmid comprising a DNA sequence which encodes a heterologous envelope protein; and
40	<i>:</i>	 a third plasmid comprising a DNA sequence of interest and HIV cis- acting sequences required for packaging, reverse transcription and integration.
	20 13.	A method of Claim 12 wherein the heterologous envelope protein is the G
15		glycoprotein of vesicular stomatitis virus (VSV G).

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10		14.	A method of Claim 12 wherein the heterologous envelope protein is the amphotropic envelope of the Moloney leukemia virus.
45		15.	A method of Claim 12 wherein the DNA sequence of interest encodes a heterologous therapeutic protein.
15	. 5	16.	A packaging cell line for producing a viral accessory protein independent
20			lentivirus-derived retroviral vector particle comprising: a) a mammalian cell; b) a first retroviral nucleotide sequence in the cell which comprises a coding sequence for a lentivirus gagpol, wherein said coding sequence
25	10		has been mutagenized to improve expression of the lentivirus gagpol proteins; c) a second retroviral nucleotide sequence in the cell which comprises the
30	15		coding sequence for a heterologous envelope protein; and d) a third retroviral nucleotide sequence in the cell which comprises a DNA sequence of interest and lentivirus cis-acting sequences required for packaging, reverse transcription and integration.
35		17.	A packaging cell line of Claim 16 wherein the heterologous envelope protein is the G glycoprotein of vesicular stomatitis virus (VSV G).
40	20	18.	A packaging cell line of Claim 16 wherein the heterologous envelope protein is the amphotropic envelope of the Moloney leukemia virus.
45		19.	A packaging cell line of Claim 16 wherein the DNA sequence of interest encodes a heterologous therapeutic protein.

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		20.	A packaging cell line comprising:
10			a) a mammalian cell;
70			b) a first retroviral nucleotide sequence in the cell which comprises a
			coding sequence for lentivirus gagpol, wherein said coding sequence has
	5		been mutagenized to improve expression of the lentivirus gagpol
15			proteins; and
			c) a second retroviral nucleotide sequence in the cell which comprises a
			DNA sequence of interest and lentivirus cis-acting sequences required
20			for packaging, reverse transcription and integration.
	10	21.	A packaging cell line of Claim 20 wherein the DNA sequence of interest
			encodes a heterologous therapeutic protein.
25			
		22.	A packaging cell line comprising:
			a) a mammalian cell;
30		•	b) a first retroviral nucleotide sequence in the cell which comprises a
	15		coding sequence for lentivirus gagpol, wherein said coding sequence ha
			been mutagenized to improve expression of the lentivirus gagpol
35			proteins; and
33			c) a second retroviral nucleotide sequence in the cell which comprises the
			coding sequence for a heterologous envelope protein.
40	20	23.	A method of producing a packaging cell line for producing a viral accessory
			protein independent lentivirus-derived retroviral vector particle, comprising co-
		•	transfecting mammalian host cells with:
45			a) a first plasmid comprising a DNA sequence which encodes lentivirus
,,			gagpol proteins, wherein said DNA sequence has been mutagenized to
	25		improve expression of the lentivirus gag and pol proteins;
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10	5		 a second plasmid comprising a DNA sequence which encodes a heterologous envelope protein; and a third plasmid comprising a DNA sequence of interest and lentivirus cis-acting sequences required for packaging, reverse transcription and—integration.
		24.	A method of Claim 23 wherein the heterologous envelope protein is the G glycoprotein of vesicular stomatitis virus (VSV G).
20		25.	A method of Claim 23 wherein the heterologous envelope protein is the amphotropic envelope of the Moloney leukemia virus.
25	10	26.	A method of Claim 23 wherein the DNA sequence of interest is a heterologous therapeutic protein.
30		27.	A method of producing a viral accessory protein independent lentivirus-derived retroviral vector particle comprising co-transfecting mammalian host cells with: a) a first plasmid comprising a DNA sequence which encodes lentivirus
35	15		gagpol proteins, wherein said DNA sequence has been mutagenized to improve expression of the lentivirus gagpol proteins; b) a second plasmid comprising a DNA sequence which encodes a
40	20		heterologous envelope protein; and c) a third plasmid comprising a DNA sequence of interest and lentivirus cis-acting sequences required for packaging, reverse transcription and integration.
45 .		28.	A method of Claim 27 wherein the heterologous envelope protein is the G glycoprotein of vesicular stomatitis virus (VSV G).

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10	29.	A method of Claim 27 wherein the heterologous envelope protein is the amphotropic envelope of the Moloney leukemia virus.
15	30.	A method of Claim 27 wherein the DNA sequence of interest encodes a heterologous therapeutic protein.
	5 31.	A viral accessory protein independent HIV-derived retroviral vector particle produced by the method comprising co-transfecting mammalian host cells with:
20		 a first plasmid comprising a DNA sequence which encodes HIV gagpol proteins, wherein said DNA sequence has been mutagenized to improve expression of the HIV gagpol proteins;
25	10	 a second plasmid comprising a DNA sequence which encodes a heterologous envelope protein; and a third plasmid comprising a DNA sequence of interest and HIV cis-
30		acting sequences required for packaging, reverse transcription and integration.
35	15 32	A method of Claim 31 wherein the heterologous envelope protein is the G glycoprotein of vesicular stomatitis virus (VSV G).
40	33	. A method of Claim 31 wherein the heterologous envelope protein is the amphotropic envelope of the Moloney leukemia virus.
	34 20	. A method of Claim 31 wherein the DNA sequence of interest encodes a heterologous therapeutic protein.
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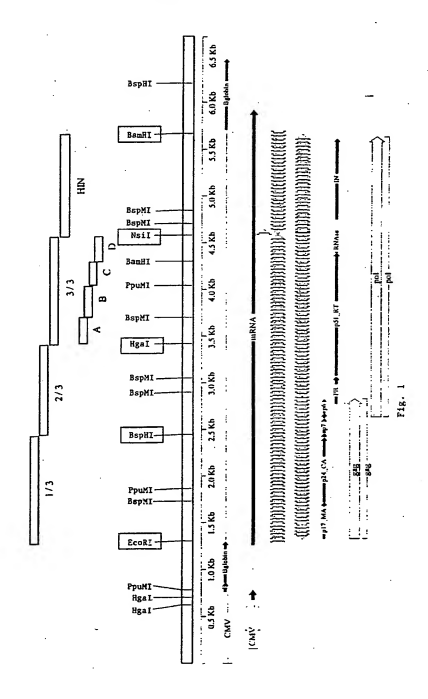
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10	35.	A viral accessory protein independent lentivirus-derived retroviral vector particle produced by the method comprising co-transfecting mammalian host cells with:
15	5	 a first plasmid comprising a DNA sequence which encodes lentivirus—gagpol proteins, wherein said DNA sequence has been mutagenized to improve expression of the lentivirus gagpol proteins; a second plasmid comprising a DNA sequence which encodes a
20		heterologous envelope protein; and a third plasmid comprising a DNA sequence of interest and lentivirus cis-acting sequences required for packaging, reverse transcription and integration.
25	36.	A method of Claim 35 wherein the heterologous envelope protein is the G glycoprotein of vesicular stomatitis virus (VSV G).
. 30	37. 15	A method of Claim 35 wherein the heterologous envelope protein is the amphotropic envelope of the Moloney leukemia virus.
35	38.	A method of Claim 35 wherein the DNA sequence of interest encodes a heterologous therapeutic protein.
40	39.	Isolated DNA encoding a codon optimized HIV gagpol.
70	40.	Isolated DNA encoding a codon optimized HIV gag.
45 ·	20 41.	Isolated DNA of Claim 40 comprising the nucleotide sequence of SEQ ID NO:4.

Isolated DNA encoding a codon optimized HIV pol.

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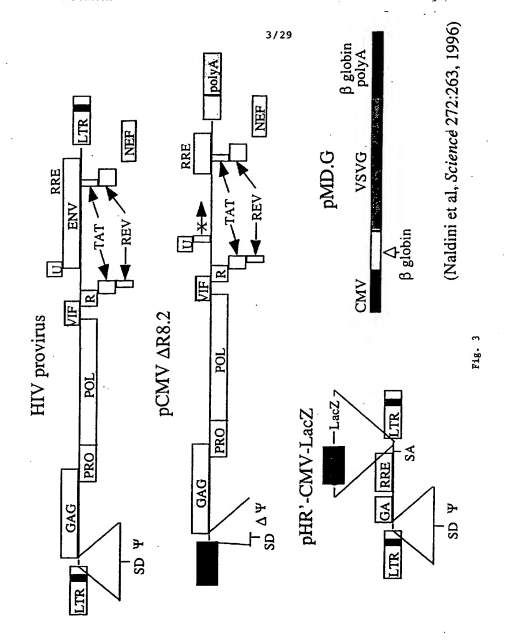
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10	43. Isolated DNA of Claim 42 comprising the nucleotide sequence of SEQ ID NO:10.
15 ·	 A method of introducing a DNA sequence of interest into a mammal comprising introducing into said mammal a viral accessory protein independent HIV- derived retroviral vector particle comprising the DNA sequence of interest.
	45. The method of Claim 44 wherein the mammal is a human.
20	The method of Claim 44 wherein the DNA sequence of interest encodes a heterologous therapeutic protein.
25	47. A method of introducing a DNA sequence of interest into a mammal comprising 10 the steps of:
30	 a) introducing into cells a viral accessory protein independent HIV-derived retroviral vector particle comprising the DNA sequence of interest; and b) returning the cells obtained in step a) to the mammal.
35	48. The method of Claim 47 wherein the mammal is a human.
40	15 49. The method of Claim 47 wherein the DNA sequence of interest is a heterologou therapeutic protein.
45	
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Codon Usage Frequencies

Amino Acid	pNL4-3	mam	Amino Acid	pNL4-3	ınanı	Amino Acid	nNI 4-3	mam
	gagpol			gagpol			promo	1110
gca Ala(A)	58	22	gga Gly(G)	55	4	cca Pro(P)	53	Ý
gcc Ala(A)	23	23	gge Gly(G)	12	20	ccc Pro(P)	2 -	2 0
gcg Ala(A)	s	11	REG Glv(G)	27	24	(1) Oct 0 Dec (1)		2 5
gcu Ala(A)	14	17	ggu Glv(G)	; v	: 2	cci Pro(F)	7 [
aga Arg(R)	63	٩	(E) (T)	, ,	3,6	ירת נוט(נ)	/7	2
(4)	3 8	2 :	כמכ הו אלה)	67	5	age Ser(S)	53	34
agg Arg(R)	٠ ج	»	can Hi s(H)	9/	21	agu Ser (S)	56	01
cga Arg(R)	4 (9	1			uca Ser (S)	. 26	'n
	0	37	ana He(I)	22	,	ucc Ser (S)	7	28
cgg Arg(R)	٣	77	and He(I)	1	, E	ucg Ser (S)	4	6
cgu Arg(R)	0	7	ann IIe(I)	26	, a	ucu Ser (S)	9	13
			(.)	2	2)
age Asin(N)	7.7	% //	cua Leu(L)	15	3	aca Thr (T)	5	1
aan Asn(N)	73	22	cric Lcu(L)	10	26		2 ==	: 4
gac Asp(D)	40	75	cug Leu(L)	=	28	ace Thr (T)		, <u>.</u>
gau Asp(D)	09	25	cun Leu(L)	=	۷.	acu Thr (T)	- 00	2.5
ngc Cys (C)	14	89	una Leu(L)	40	2	IIO Trn(W)	1001	. 6
ugu Cys (C)	92	32	ung Leu(L)	13	9		3	3
			aaa Lys (K)	69	×	uac Tyr (V)	35	7
caa Glu(Q)	95	12	aag Lvs (K)	-	£ 6	near Ty= (Y)	9 7	<u> </u>
cag Gln(Q)	44	88	(-)-(-0-		70	(i) if nen	4/	97
			ang Met (M)	90	100	gua Val (V)	58	2
(1)	Q.F	,				guc Val (V)	13	25
gaa Giu(E)	2 ;	?	unc Phe (F)	- 40	08	gug Val (V)	91	64
gag Oin(E)	30	75	unn Phe (F)	09	20	guu Val (V)	4	7

Fig. 2



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- Regulates HIV gene expression by promoting cytoplasmic levels of unspliced and singly spliced mRNAs
- Postulated to affect splicing, stability, transport, and translation

Fig. 4

Codon Optimization of HIV gagpol

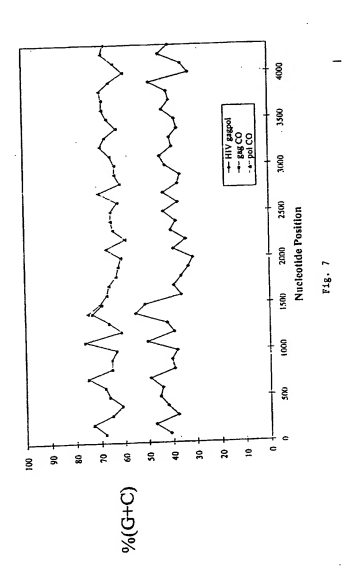
- Remove A-rich instability elements
- Improve translational efficiency
- Reduce risk of recombination with transfer vector

Inactivation of Inhibitory Sequences in gag Schwartz, S., et al.

tta gac aag ata gag gaa g<u>ag caa aac aaa agt aag aaa aaa gca cag caa gca gca gc</u>t atg ggt gcg aga gcg tca gta tta agc ggg gga gaa tta gat cga tgg gaa aaa att cgg tta agg cca ggg gga aag aaa aaa tat aaa tta aaa cat ata gta tgg gca agc agg gag <u>aca gta gca acc ctc tat tgt gtg cat caa agg ata gag ata aaa gac acc aag gaa gct</u> cta gaa cga ttc gca gtt aat cct ggc ctg tta gaa aca tca gaa ggc tgt aga caa ata ctg gga cag cta caa cca tcc ctt cag aca gga tca gaa gaa ctt aga tca tta tat aat G G SCC M2 gac aca gga cac agc aat cag gtc agc caa aat tac 35 3 5 5 5 CGC

Fig. 6

Nucleotide Content of HIV gagpol



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Alignment Report of Codon optimization (gag).MEG, using Clustal method with PAM250 residue weight table.

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792	M						v	L	5					D		NL4-3 genbank.SEG
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		840										870				-
837	W	E	ĸ	I	R	L	R	P	G	G	K	ĸ		Y	K	NL4-3 genbank.SEQ
837						S TTA										
1364 1364		E GAG	X AAC	I AT	R CGC	L CTG	R	P CCC	G GGG	G GG	K S AA	K B AA		Υ 14 T =	. AAC	pHDMHgpm2.seq
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882	L	K	H	I	v	W	À	5	R	Ε	L	E	R	F	A	NL4-3 genbank.SEQ
882 1409		L AAJ	CA1	: AT/	i GTZ V	I TGG	GCA A	AGC 5	: AGC R	GA(CT)	E GAV	A CG2 R			
1409						TGG								F TTC	A GCC	pHDMHgpm2.seq
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927 1454	CL.	'AAT N	, CCI	GGC	CTI L	TTA L	GAG E	ACA T	. TCA S	GA! E	GGC					
1454						CIG						C TGC	R CGC	Q CAG	I ATC	pHDMHgpm2.seq
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972 972	L CTG	G GGA	Ç CAG	L	Q CAA	P CCA	5 TCC	L	Q CAG	T	G GGD	S TC2	E	E CAA	L	NL4-3 genbank.SEQ
1499	L	G	Q	L	Q	P	s	L	Q	T	G	5	2	E	L	pHDMHgpm2.seq
1499	CTG	GGC	CAG	CTG	CAG	CCC	TÇC	CTG	CAA	ACC	GGC	TCC	GAG	GAG	CTG	
		.020										050				•
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1017		-				ACA				_						NL4-3 genbank.SEQ
1544	R	s	L	¥	N	T	I	A	V	L	Y	c	v	Ħ	Q	pHDMHgpm2.seq
1544	CGC	TCC	CTG	TAC	AAC	ACC	ATC	GCC	GTG	CTG	TAC	TGC	GTG	CAC	CAG	
						1	080									
1062	R	I	D	v	ĸ	Đ	T	ĸ	E	A	L	D	K	I	E	NL4-3 genbank.SEQ
1062						GAC										
1589 1589	R	I	GAC	CTC.	K	Cac D	T	K	E E	A GCC	CTG	ESC.	K	I	E	pHDMHgpm2.seq
				7.7				~~	47.0		-:0	- GAL	7110	AIC	wAG.	
	1	110									1	140				
1107	E	Ε	Q	N	ĸ	S	ĸ	K	К	Α	Q	Q	A	A	A	NL4-3 genbank.SEQ
1107 1634	GAA			aac N	AAA.	AGT .	AAG. K									- times
		ಕ ಎ	Q CAG			TCC .		K Aag	k aag	A GCC	CY.C.	Q CAG	A GCC	GCC	A GCC	pHDMHgpm2.seq
										~	٠.٠		300			

Fig. 8A

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Alignment Report of Codon optimization (gag).MEG, using Clustal method with PAM250 residue weight table.

																,
						1	170									
1152	D	T	G	N	N	s	Q	٧	3	Q	М	¥	P	I	٧	NL4-3 genbank.SEQ
1152				AAC N	AAC N	AGC S	CAG	GTC	AGC S	CAA Q	AAT	Y	P	ATA	GTG V	pHDMHqpm2.seq
1679 1679	D D	T ACC	GGC	AAC	AAC	TCC	CAG									piibi siqpiib i seq
10.,																
		200	-		_						1	230				
1197	Q	N	L	Q	G	Q	M	٧	H	Q	A	I	5	P	R	NL4-3 genbank.SEQ
1197					GGG G	CAA	ATG M	GTA V	CAT H	CAG Q	GCC	ATA I	TCA	CCT	AGA R	pHDMHgpm2.seq
1724 1724	Q CAG	N AAC	L	Q CAG	GGC	CAG	ATG	GTG	CAC	CAG						Printing Printing
							T									•
						1	260							<u>. </u>		
1242 1242	T	L	N	A	W	V	K	V	V	CAA	E E	K NAC	A CCT	F		NL4-3 genbank.SEQ
1242	ACT T	TTA L	AAT N	GCA A	TGG W	GTA V	AAA K	V	V SIA	E	E	R	A	F	3	pHDMHgpm2.seq
1769	ACC	CTG				GTG		GTG	GTG	GAG	GAG	AAG	GCC	TTC	TCC	
		-					-									•
	1	290									1	320				•
1297	P	E	V	I	P	M	F	S	A	L	S	E	G	A	T	NL4-3 genbank.5EQ
1287		GAA E	GTA V	ATA I	CCC	ATG M	TTT	TCA S	GCA A	L	5	E	G	A	T	pHDMHgpm2.seq
1814	CCC	GAA	GTC	ATC												,
							_									-
							350									
1332 1332	P	Q	D	L	N	T	M	L	N	T	V GTG	G	G	B CAT	C22	NL4-3 genbank.SEQ
1859	P	0	D	L	N	T.	M	<u>.</u>	N	Ţ	v	G	G	Ħ	Q	pHDMHgpm2.seq
1859	ccc	CĀG	GAC	CTG	AAC	ACC	ATG	CTG	AAC	ACC	GTG	GGC	GGC	CAC	CAG	
	_	-														•
	1	380										410				
1377	A	Α	M	ÇAA	M	L	K	E	T	I	N	E	E	A	A	NL4-3 genbank.SEQ
1377	GCA A	GCC	ATG M	CAA Q	ATG M	TTA L	X	E	T	I	N	E	E	A	A	pHDMHgpm2.seq
1904	GCC	ecc	ATG	CAG	ATG	CIG	AAG	GAG	ACC	ATC	AAC	GAG	GAG	GCC	GCC	
							_									-
						1	440									•
1422	Z	W	D	R	L	H	P	٧	H	A	G	P	I	A A	CC A	NL4-3 genbank.SEQ
1422	GAA E	TGG	GAT D	AGA R	TTG L	CAT	CCA P	GTG V	H	A	GGG	P	Ĭ.	A	P	pHDMHgpm2.seq
1949	GAG	TGG	GAC	CGC	CTG	CAC	CCC	GTG	CAC	GCC						
																•
	_ 1	470						•				500				•10
1467	G	Q	M	R	E	5	R	G	5	D	I	A	G	T	T	NL4-3 genbank.SEQ
1467				AGA	GAA E	CCA P	AGG R	GGA G	AGT S	eyc eyc	ATA I	A	GGA G	ACT	T	pHDMEgpm2.seq
1994 1994	G	C3G	M ATG	R CGC	GAG	CCC	CGC									*
1774	300		~**	505												

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Alimment Report of Codon optimization (gag) MEG, using Clustal method with PAM250 residue weight table.

-ng,	0,14 . 40	p u		-, op.		ψ., _{(G} .	3,	,								
						1	530									
1512	s	T	L	Q	E	Q	I	G	W		T	H	N			NL4-3 genbank.SEQ
1512	AGT	ACC	CTT	CAG	GAA	CAA	ATA	GGA	TGG	ATG	ACA	CAT	AAT	CCA	CCT	
2039	S	T	L	Q	E	Q	I	G	W	M	T	H	N	P	P	pHDMHgpm2.seq
039	TCC	ACC	CTG	CAA	GAG	CAG	ATC	GGC	TGG	ATG	ACC	CAC	AAC	CCC	ccc	
		-														· _
		560										590				
.557	1	2	V	G	E	I	Y	K	R	W	I	I	L	G	L	
	ATC				E	I	Y	K	R	M. 100	I	I	L	G	L	pHDMHqpm2.seq
084	I ATC	CCC	V GTG	G GGC												buningham . ned
.004																
						1	620									
602	N	К	I	7	R	M	Y	5	₽	T	5	I	L	D	I	
602	aat	AAA	ATA													
2129	N	K	I	v	R	M	Y	5	P	Ť	5	I	L	D		pHDMHgpm2.seq
2129	AAC	AAG	ATC	GTG	CGC	ATG	TAC	. TCC	CCC	ACC	TCC	ATC	CrG	GAC	ATC	
	1	650									1	680				•
647	R	Q	G	P	K	ε	P	ε	R	D	Y	v	D			NL4-3 genbank.SE(
	AGA															-117161
2174	R	Q	G	P	X AAG	E	P	F	R	D	Y	V	D CAC	R		pHDMHgpm2.seq
2174	CGC	٠,			AAG	GAG		110		<u> </u>			W.C			
						1	710									
692	Y	К	T	L	R	A	E	Q	A	S	Q	E	V	K		NL4-3 genbank.SEQ
1692	TAT	aaa	act	CTA												
2219	Y	K	T	L	R	A	E	Q	A	5	Ç.	E	V CTA	K	N	pHDMMgpm2.seq
2219	TAC	AAG	ACC	CTG	CGC	GCC	GAG	CAG	GCC	TCC	LAG	GAG	GIA	MAG	AAC	
	1	740									1	770				
737	W	м	T	Е	T	L	L	٧	Q	N	A	N	P	D	С	NL4-3 genbank.SEQ
737	TGG	ATG	ACA	gaa									CCA	GAT	TGT	
2264	W	M	T	E	T	L	L	٧	Q	N	A	N	P	D		pHDMHgpm2.seq
2264	TGG	ATG	ACC	GAG	ACC	CTG	CTG	GTG	CAG	AAC	GCC	AAC	ccc	GAC	TGC	
						1	800									
782	ĸ	T	I	L	ĸ	A	L	G	P	G	A	T	L	Е	E	
782	AAG	ACT							CCA		GCG	ACA	CTA	GAA	GAA	
309	K	T	I	L	K	A	L	G	5	G	A	T	i	E	E	pHDMHgpm2.seq
309	AAG	ACC	ATC	CTG	AAG	GCC	CTG	GGC	CCC	GGC	GCC	ACC	CTG	GAG	GAG	
	1	830									. 1	860				0 =-
827	M	м	T	A	c	Q	G	v	G	G	Р	G	H	К		NL4-3 genbank.SEQ
827	ATG	ATG	ACA	GCA	TGT	CAG	GGA	GTG	GGG	GGA	CCC	GGC	CAT	AAA	GCA	
2354	М	M	T	A	С	Q	G	v	٠G	G	P	G	H	ĸ	A	pHDMHgpm2.seq
2354	ATG	ATG	ACC	GCC	TGC	CAG	GGC	GTG	GGC	GGC	CCC	GGC	CAC	AAG	GCC	

Fig. 8C

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Alignment Report of Codon optimization (gag).MEG, using Clustal method with PAM250 residue weight table.

							890									-
1070		v		A	E		M	s	Q		T	N	Р	A	Ŧ	NL4-3 genbank.SEQ
1872 1872			L													WIN-3 GEIMHINION
2399	R	v	L	A	E	A	M	5	Q	٧	T	N	P	A	Ť	pHDMHgpm2.seq
2399					GAG	GCC	ATG	TCC	CĀĀ	GTC	ACC	AAC	CCC	GCC		
		920									1	950				•
						_	-:-		_	N		R	ж	T	v	NL4-3 genbank.SEC
1917 1917	I	M	I.		K	G	N aat	F	R		Car					ML4-3 genbank.322
2444	I	M	I	Q	K	G	N.	F	R	N	Q.	3	ж	7	v	pHDMHgpm2.seq
2444																•
							980									
1962	K	c	2	N	c	G	K	E	G	H	I	A	X Dan	N	C TCC	NL4-3 genbank.SEQ
1962			F	AAT N	C	GGC	X	E	G	H	I	A	K	Ň	C	pHDMHqpm2.seg
2489 2489	X AAG	TGC														huntardhum 1004
2403	7,7,0															
	2	010									2	040				
2007	R	A	P	R	ĸ	ĸ	G	С	M	K	С	G	К	Ξ	G	NL4-3 genbank.SEQ
2007	AGG	GCC	CCT	AGG	AAA											
2534	R	A'	P	R	K	ĸ	G	С	W	ĸ		G	X	Ξ		pHDMHgpm2.seq
2534	CGC	GCC	ccc	CGC	AAG	AAG	GGC	TGC	TGG	AAG	TGC	GGC	MAG	GNG	GGC	
					-	· 2	070									
2052	H	0	м	3	D	С	' T	E	R	Q	A	Ж	F	L	G	NL4-3 genbank.SEQ
2052		CAA	ATG		GAT	TGT	ACT	GAG	AGA	CAG	GCT	aat	TIT	TTA	GGG	
2579	H	Q	M	K	D	С	T	Ξ	R	Õ	Α	N	F	5	G	pHDMHqpm2.seq
2579	CAC	CAG	atg	AAA	GAT	TGT	ACT	GAG	AGA	CAG	GCT	AAT	TIT	TTA	GGG	
		100									2	130				
					s	н	х	G	Я	P	G	N	F	=	Q	NL4-3 genbank.SEQ
2097 2097	K	I	TGG	P									TTT			not 5 genoum
2624	R	I	W	P	s	н	К	G	R	P	G	N	F	L	Q	pHDMHgpm2.seg
2624	AAG	ATC	TGG	CCT	TCC	CAC	AAG	GGA	AGG	CCA	GGG	aat	TTT	CTT	CAG	
							160									
					_		160	P	P	E	E	s	E	R	e	NL4-3 genbank.SEQ
2142 2142	5	R	6	E	P.	T	A									nt4-3 deimank.255
2142	AGC S	AGA R	P	E E	P	T	A	P	P	E	ε	3	£	R	F	pHDMEgpm2.seq
2669	AGC	AGA	CCA	GAG	CCA	ACA	GCC	CCA								,
••••																
		190.									2	220				
2197	G	E	E	Ŧ	T	T	P	S	Q	K	Q	Ξ	5	I		NL4-3 genbank.SEQ
2197		GλA									CAG	GAG	CCC	ATA	GAC	
2714	G	E	E	T	T	T	P	\$	Q	K	C3G	E 636	P	I 272		pHDMHgpm2.seq
2714	GGG	CYY	GAG	ACA	ACA	ACT	CCC	rc:	فالات	MAG	فناحب	GAG.	LUG	A.A	unc	

Fig. 8D

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Alignment Report of Codon optimization (gag) MEG, using Clustal method with PAM250 residue weight table.

						2	250									
2232	ĸ	Ė	L	Y	P	L	A	3	L	R	3	L	F	G	s	NL4-3 genbank.SEQ
2232	AAG	GAA	CTG	TAT	CCL	TTA	GCT	TCC	CTC	AGA	TCA	CTC	TTT	GGC	AGC	
2759	ĸ	E	Ł	Y	P	L	A	3	L	R	S	L	F	G	S	pHDMHgpm2.seq
2759	AAG	GAA	CTG	TAT	CCT	TTA	GCT	TCC	CTC	AGA	TCA	CTC	TTT	GGC	AGC	•
																
	2	280														
277	D	ъ	s	5	Q	Į.,										NL4-3 genbank.SEQ
277	GAC	CCC	TCG	TCA	CAA	TAA										
804	D	P	5	5	Q.											pHDMHgpm2.seq
2804	CAC	ccc	TCG	TCA	CAA	TAA										

Fig. 8E

Alignment Report of Codon Optimization (pol) MEG, using Clustal method with PAM250 residue weight table.

												_				· · · · · · · · · · · · · · · · · · ·
	21	390			•						2	120				
2087	F	F	R	E	D	L	A	2	P	Q	G	X	A	R	E	NL4-3 genbank.SEQ
2087	TTT	TTT	AGG	GAA	GAT	CTG	GCC	TTC	CCA	CAA	GGG	AAG	GCC	AGG	GAA	
2085	E*		R	F.	מ	L.	А	F	P	Q	G	K	А	ĸ	E	pNL4-3.seq
2085								TTC	CCA	CAA Q	G	K	A	R	E	pHDMHgpm2.seq
2612 2612	F	F	R	E	D	L	A	E TTC	CC.3							bantmahmma
2612	TTT	TTT	AGG	GAA	GAI	CIG	GCC	110				•••				
		<u></u> -					150									•
				Ė	0		R	Α.	N	5	2	T	R	R	Ε	NL4-3 genbank.SEQ
2132 2132	E	S TCT	5 TC2	GAG	CAG	ACC	AGA	ecc.	AAC	AGC	ccc	ACC			GAG	
2120	~	•	S	F.	0	7	9	A	N	5	2	T	ĸ	к	E.	pNL4-3.seq
2130	TTT	TCT	TCA	GAG	CAG	ACC	AGA	GCC	AAC	AGC	CCC	ACC	AGA	AGA	GAG	
	-		•	7	0	т	2	A	N	S	2	T	ĸ	×	2	phrmidbur.sed
2657	TTT	TCT	TCA	GAG	CAG	ACC	AGA	GCC	AAC	AGC	CCC	ACC	AGA	Alari	GAG	
		-									2	210				•
	2	180									L	s	E	A		NL4-3 genbank.SEQ
2177	L CTT	Q	V	W	G	R	D	N	N	S TCC						
				TGG	GGA	AGA R	D	N	N	S	L	s	E	A	G	pNL4-3.seq
2175	L CTT	CAG	CTT.	TGG	GGA	AGA	GAC	AAC	AAC	TCC	CTC	TCA	GAλ	GCA	GGA	
		_	1/	T.J	<u> </u>	2	D	N	N	5	L	Þ	5	^	G	buttendburg. acd
2702	CIT	CĀG	GTT	TGG	GGA	AGA	GAC	AAC	AAC	TCC	CTC	TCA	GAA	GCA	GGA	
							1									-
							240				_	5	Q	ī	Ŧ	NL4-3 genbank.SEQ
2222	A GCC	D	R	Q	G	T	V	3	ىسلىس <u>د</u>	5 360	AAC E					
		GAT D	AGA R	Q	G	T	A A	S	F	5	F	P	Q	I	T	pNL4-3.seq
2220	A GCC	ע פאד	PCP V	CAA	GGA	ACT	GTA	TCC	TTT	AGC	TTC	CCT	CAG	ATC	ACT	
2717		-		0	G	т	v	S	F	5	F	P	v	•	1	puning puz. sed
2747	GCC	GAT	AGA	CÄA	GGA	ACT	GTA	TCC	TTI	AGC	TTC	CCT	CAG	ATC	ACT	
												1				-
	2	270									2	2,00				
2267	L	W	Q	R	P	L	V	T	I	K	I	G	G	Q		N14-3 genbank.SEC
2267	CTT	TGG	CAG	CGA	ccc		GTC	ACA	ATA	AAG	ATA	GGG	GGG	CAA Q	. TTA L	pNL4-3.seq
2265	L	M	Q	R	p	L	V	T	I	K	I ata	CCC.				
	CII				. CCC	CTC L	GTC	ACA T	I	K	I	G	G	Q	L	pHDMHgpm2.seq
2792	CTT	W	C)C	R	CCC	CLC P	GTC	ACA	ATA	AAG						
2792	CIT	166	unu	Cun				,,,,,								_
						2	330									
2312		E	A	L	L	D		G	A	Đ	D	T	v	L		NL4-3 genbank.SEQ
2312	AAG	GAA	GCT	CTA	TIA	GAT	ACA	GGA	GCA	GAT	GAT	ACA	GTA	TIA	(GYY	
						D	7	G	A	D	υ	-	٧	-		2MT4-2.26d
		E	A	L	L		-									
		E GAA	A GCT	CTA	TTA	GAT	ACA	GGA	GCA	GAT	GYI	ACA	GTA	. TTA	. GAA	nUDMicrom? sec
2310 2310	K Aag	GAA	GCT	CTA	TTA	GAT	ACA T	G		U	ν	·	•		-	Subiarabire

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Alignment Report of Codon Optimization (pol).MEG, using Clustal method with PAM250 residue weight table.

					-											
		236)									2390)			
235	_	•	-				R	W	K	P	K		1	G	G	NL4-3 genbank.S
2357			G A	T TI	e cc	A GG	A AG	A TG	S AA	A CC	AA A	TA A	G AT	A GG	G GG	Α
2355	_	M		_			R	W	ĸ	P	K	M	I	G	G	pNL4-3.seg
2355					G CC						AA A	A AT	G AT	A GGG	GG	Α .
2882		. M				G	R	W	K	P	ĸ	М	I	G	G	pHDMHgpm2.seq
2882	GA	3 AT	G AA	C CT	G CC	C GG	CGC	TGG	S AAG	S CCC	: AA	G AT	3 ATC	GGG	GGG	•
							2420									-
402		G	G			ĸ	v	7 G		Y	D	Q	I	L	I	NL4-3 genbank.SE
2402	AT	GG	A GG		T AT						GA1	CAC	ATA	CTC	AT/	
2400	_	G	G	_		K	v	R	Q	Y	D	Q	I	L	I	pNL4-3.seg
2400			A GG		T AT							CAC	ATA	CTC	: ATA	`
2927		G.	G 		I	X	v	R	Q	Y	D	Q	I	L	I	pHDMHgpm2.seq
921	ATC	. 66	. 66	C Tr	C AT	; AAA	GTC	CGC	CAG	TAC	GAC	CAC	ATC	CTG	ATC	
		2450										2480				-
447	E	I	C	G	Н	ĸ	A	I	G	T	٧	L	٧	G	P	NL4-3 genbank.SE
2447	GAA	ATO	TG		A CAT								GTA	GGA	CCT	
445		I	C	G	H	K	A	I	G	T	V	L	v	G	P	pNL4-3.seq
445 972					CAT											
	_	I DTC	C	G	H	K	A	I	G	T	V	L	V	G	P	pHDMHgpm2.seq
312			. 10	- 660	CAC	MAG	GCC	AIC	GGC	ACC	Gre	CTG	GTG	GGC	ccc	
						2	510									
492	T	P	γ	N	I	I	G	R	N	L	L	T	Q	ī	G	NL4-3 genbank.SE
492					: ATA											
490	T	P	V	N	I	1	G	R	N	L	L	Ť	Q	I	G	pNL4-3.seq
017	T	P	V	. AAL N	ATA											
					I DTC	I	G	R	N	L	L	T	Q	Ţ	G	pEDMHgpm2.seq
01,	ACC		G16	MAL	ATC	AIC	GGC	CGC	AAC	CTG	CrG	ACC	CAG	ATC	GGC	
	2	540									2	570				•
537	С	T	L	N	F	6	1	S	P	I	E	T	٧	P	v	NL4-3 genbank.SEG
537		ACT		AAT		CCC						ACT	GTA	CCA	GTA	
535	С	T	L	N	F	P	I	S	P	I	E	T	v	P	V	pNL4-3.seq
535				AAT		CCC								CCA	GTA	
062	С	T	L	N	F	P	I	s	P	1	E	T	v	P	٧	pHDMHgpm2.seq
62	TGC	ACC	CTG	AAC	TTC	ccc	ATC	TCC	CCC	ATC	GAG	ACC	GTG	CCC	GTG	
				***		2	500								_	
82	ĸ	L	K	P	G	м	D	G	P	x	v	K	Q	W	P	NL4-3 genbank.SEC
82	AAA	TTA	AAG	CCA	GGA	ATG	GAT	GGC					CĀA ·			germann. 304
80	K	I.	ĸ	P	G	M	D	G	P	K	v	ĸ	Ω	W	P	pNL4-3.seq
80	AAA	TTA	AAG	CCA	GGA	ATG	GAT	GGC	CCA .	AAA	GTT					
30																
07	K	L	ĸ	P	G	M	D	G	P	ĸ	v	K	Q	W	P	pHDMHgpm2.seg

Fig. 9B

Alignment Report of Codon Optimization (pol).MEG, using Clustal method with PAM250 residue weight table.

																•
		263	0									266	0			
262	7 T	•		Ξ	2 2	7	K	A	L	v	, E	_		T	Ε	
262				_							_				A GA	
262					E 8									T		-
252					A AF	TA AJ	A AA	A GC	A TT	A GT	A GA	a at	T TG	r ac	A GA	
315					2 2					_ v				T		pHDMHgpm2.seq
315.	z Cr	G AC	:C G	AG G	AG AA	IG AT	CAA	G GC	CCT	G GT	G GA	G AT	C TG	CAC	C GAG	;
							2690)				_				-
267		_		¢ 1	-		_	s	K	ī	G			N	P	
267							A AT		A AA		r GG	G CC	T GA	AA.	r cc	\ .
2670							I	S	K	I	G	P	E	N	P	pNL4-3.seq
2570 3191															r cca	
				C E			I TATO	S TC:	K - 220	I Tate	G Car	P	E - C30	N 334	P CCC	pHDMHgpm2.seq
						- ,							- Gra	, ,,,,,		•
		27,20)									2750				-
2717						-	λ	I	К	ж	К	Đ	s	T	ĸ	
2717														ACI	. AAA	
2715	-	N	_			_	. A	I	К	K	X	D		T	K	pNL4-3.seg
2715 3242		aa: N	r ac T			a TT	GCC A	: ATA I	AAG X	AA. K					· AAA	
	-										K AAC	ם זבה:	 	T	K : AAG	pHDMHgpm2.seq
							2780									
2762		R	К		V	D	F	R	Ε	L	И	X	R	T	Q	NL4-3 genbank.SEQ
2762					A GT											
2760 2760	W TGC	R	X 		V.	D	F	R	E	L	N	K	. R	T	Q	pNL4-3.seq
287	N CC	, AG. R	۰۰۰۰ K		A GT.	י מעז	F	R	E	L	W	K	3	ACT	Q	pHDMHqpm2.seq
297					GTG	_	_									parangpaz.seq
		2810									- 2	94C				
807	D	F	W	E	٧	Q	L	G	I	P	. H	5	Α	G	I,	NL4-3 genbank.SEQ
807	GAT				GTT								GCX			
805	D GAT	F	W TG:	E Car	V LGTT	. caa	L	ees G	I	CC3	H Car	5 5	A GC3	G	L	pNL4-3.sec
332	ם D	F	W	E	. G.	~	L	G	I	P	ä	Ρ.	A	G	L	pHDMHqpm2.seq
					GTG	-								_		Sun mid him o a e d
						2	870									
952	K	Ç	K	K	S	٧	7	٧	L	D	٧	G	D	A	Y	NL4-3 gembank.SEC
852					TCA											
850	K	Q	K	K	S	V	T	V CTD	L	D	V	G	D	A	Y.	pNL4-3.seq
850 377	K	Q	AAA X	. aaa K	TCA S	GTA V	ACA T	GTA V	L	D	V	GGC	D GAT	GCA A	TAT Y	numumn? eng
377					TCC											pHDMHgpm2.seq
			0	~~0					~ . ~		J. J				•~~	

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Alignment Report of Codon Optimization (pol).MEG, using Clustal method with PAM250 residue weight table.

		-														
		2900										2930				
2897	F	3	v	P	L	D	К	D	F	R	ĸ	<u> Y</u>	T	A	F	 NL4-3 genbank.S
2897	TTT			ccc		GAT							' AC	, ech	TT	7
2895	-	S	. V	P	L	ם	K	D	F	R	K	Y	T	A	F	pNL4-3.seq
2895					TTA											•
3422	-		v	. eee	L	D	K	D	F	R	K	Y	T	A	F	pHDMHgpm2.seq
3422	TTC	. TCC	. Gre	, (:.	CTG	GAC	AAG	GAC	Tre	CGC	AAG	TAC	ACC	GCC	TTC	
	_		-				960									-
2942			P	s	I	— <u>.</u>	N	Ε.	T	P	G	r	R	Y	Q	- NL4-3 genbank.S
2942	ACC	_	_	AGT		AAC						_			CAG	
2940	T	I	P	5	I	N	N	E	T	Ъ	G	I	R	Y	Q	pNL4-3.seq
2940	ACC	ATA:	CCT	AGT	ATA	AAC	AAT	GAG	ACA	CCA						i pros stacq
3467	T	I	P	S	I	N	N	E	T	P	G	1	R	Y	Q	pHDMHgpm2.seq
3467	ACC	ATC	CCC	TCC	ATC	AAC	AAC	GAG	ACC	ccc	GGC	ATC	CGC	TAC	CAG	
		2990										3020				_
2987	- <u>-</u>	N		L	P	Q	G	W	К	G						
2987	-				CCA						S	P	A	I	F	NL4-3 genbank.S
2985	Y	N.	V	L	P	Q	G	M	K	G	S	P	A	I		
2985	_		-		CCA										THTC E	pNL4-3.seq
3512	Y	N	v	L	P	Q	G	W	K	G	s	P	A	I	. IIC	pHDMHqpm2.seq
3512			GTG	_	CCC											
																_
						3	050					_				
032	Q	С	S	M	T	K	I	L	E	P	E	R	К	Q	И	NL4-3 genbank.Si
3032					ACA				GAG					CAA	aat	
1030	Q	С	S	M	T	K	I	L	E	₽	F	R	K	Q	N	pNL4-3.seq
030					ACA				GAG				AAA			
557	Q	C	S	M	T	K	I	L	Ξ	P	F	R	K	Q	N	pHDMHgpm2.seq
331	CAG	TGC	TCC	ATG	ACC	AAG	ATC	CTG	GAG	CCC	TTC	CGC	AAG	CAG	AAC	
		080									3	110				
077	P		I	v	I	Y	Q	Y	M	D	D	L	Y	v	G	NL4-3 genbank.SE
077	CCA			GTC	ATC		CAA				-					o germank.be
075	P	D	1	v	I	Y	Q	Y	M	D	D	L	Y	v	G	pNL4-3.seq
075	CCA	GAC	ATA	GTC	ATC	TAT	CAA	TAC	ATG	GAT	GAT	TTG	TAT	GTA	GGA	•
602	P	D	I	V	I	Y	Q	Y	M	D	D	L	Y	V	G	pHDMHgpm2.seg
602	ccc	GAC	ATC	GTG	ATC	TAC	CAG	TAC	ATG	GAC	GAÇ	CTG	TAC	GTG	GGC	
						21	40							·		
122	s	D	L	E	<u> </u>	G 3,	ــــــــــــــــــــــــــــــــــــــ	н	R	T	K	ī	Ε	E	.	
	TCT	-	_	_	ATA	-	Q CAG			_			GAG	_	L	NL4-3 genbank.SE
120	S	D	L	E	I	G	Q	E CYT	R	T	K	I	E	eaa E		m)(7.4. 2
	-	-	_	_	ATA :							_			L CTG	pNL4-3.seq
120	TUT															
120 647	S	D	L	E	I	G	Q	Ħ	R	T	K	I	E	E	L	pHDMHqpm2.seq

Fig. 9D

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Alignment Report of Codon Optimization (pol) MEG, using Clustal method with PAM250 residue weight table.

		31	70							_			32	00				
316		₹.		Н	L	L	R	OF			7	,	F	P	D	К	K	NL4-3 genbank
316	7 AC	A C	AA (AT	CTG	TTO	AG(TG:	G GG	A TI	T AC	C A	CA C	CA	GAC	AA	AA A	A
316	5 F	₹ .	Q	H	L	L	R	W	G	E	' 1		r	P	D	×	v	-177 4 3
316	5 AG	ia c	AA C	AT	CTG	TTG	AGG			A TT			CAC	CA	GAC	AAJ	AA A	Α
369			Ω.	H	L	L	R	W	G	F	. 1	1	r	P	D	K	K	pHDMHgpm2.seq
309	2 CG	ec c	AG (AC	CTG	CTG	CGC	TGO	G GG	C TT	C AC	CAC	:c c	CC	GAC	AAC	AA .	G
				_			,	3230	***			_			_			_
321			-	K	E	P	P	F	L	W	M	- 0	;	Y	E	L	н	- NL4-3 genbank.
321						CCI	CCA	TTC	CT	r TG	G AT	G GC	T T	AT	GAA	CTC	CA	r
321				K	E	P	P	F	L	W	M	G	;	Y	F.	t.	12	mNT 4 2 ac-
3210	CA	TC	IG A	AA :	GAA	CCI	CCA	TTC				G GG	T T	AT	gaa	CTC	CAT	
373				K	E	P	P	F	L	v	M	G	,	Y	Ε	L	H	pHDMHgpm2.seq
3/3	7 CA		G A	AG (GAG	CCC	ccc	TTC	CTC	TG	G AT	GGG	C T	AC (GAG	CTG	CAC	
		326	0										329	0				_
3257				K	W	T	V	Q	P	I	v	L	1	,	E	к	D	- NL4-3 genbank.S
3257	CC	L CY	TA	AA :	rgg	ACA	GTA	CAG	CCI	AT.	GT(CT	G C	a c	SAA	AAG	GAC	yemanx
3255	P	D	, ,	K	W	T	v	0	P	I	v	L		>	E	W	D	-NT 4 2
3255	CC	r ga	TA	VA 2	PGG	ACA	GTA	CAG	CCT	ATA	GT	CT	G CC	:A (AA	AAG	GAC	i i i i i i i i i i i i i i i i i i i
3782		D		ζ	W	T	V	Q	₽	1	v	L	I	•	E	K	D	pHDMHgpm2.seq
3/82	CCC	: GA	C A	iG 1	."GG	ACC	GTG	CAG	CCC	ATC	GTG	CT	G CC	:C 6	AG	AAG	GAC	
							3	320							_	—-		•
3302		W	7		٧	N	D	'I	Q	ĸ	L	v	G	_	ĸ	L	N	- NL4-3 genbank.S
3302	AGC	TG	G AC	T G	TC .	AAT	GAC	ATA	CAG	AAA	. TTA	GTO	GG	A A	AA ·	TTG	AAT	MD4-3 Genbank.3
3300	\$	W	1	•	V	N	D	I	Q	K	L	v	· G		K	L	N	pNL4-3.sec
3300	AGC	TG	3 AC	T G	TC 2	aat	GAC	ATA	CAG	AAA	TTA	GTG	GG	АА	AA '	TTG	AAT	huns asped
3927	ន	W	T	,	v	N	D	1	Q	K	L	v	G		ĸ	T.	N	pHDMHgpm2.seg
827	TCC	TG	S AC	CG	TG A	AAC	GAC	ATC	CAG	AAG	CTG	GTG	GG	C A	AG (CTG	AAC	Prostangpine . 3 c g
		350											3380	,				
347	W	A	s		Q	I	Y	A	G	ī	К	v	R		2	L	С	NY 4-2 combacts of
3347	TGG	GC	. AG	rc	AG 2	ATT '	TAT	GCA								TA '		NL4-3 genbank.SI
3345	₩	A	S	•	Q	I	Y	A	G	I	K	v	R		2	t.	_	pNL4-3.seg
345	TGG	GCA	AG	rc	ag a	TT!	TAT	GCA	GGG	ATT	AAA	GTA	AG	s ci	AA 1	TA '	IGT	
872	74	A	5	ç	Q	I	Y	A	G	I	K	v	R		•	T.	_	pHDMHgpm2.seq
872	TGG	GCC	TC	: C/	AG A	ATC :	TAC (SCC	GGC	ATC	AAA	GTC	CGG	C	AG C	TG :	rgc	·
							34	10										
392	К	L	L	-,9	·	G	T	K	A	L	Ŧ	E	v	v	,	P	L	N7.4 3 mambant ==
392	AAA	CTT	CTT				CC A			CTA	ACA	GĀA	GTA	GT	'A C	ca c	ברד.	NL4-3 genbank.SE
390	K	L	L	P	₹ '	G	T	K	A	L	T	E	V	v	,	P	T.	pNL4-3.seq
390	AAA	CTT	CTI	' AG	G G	GA A	CC A				ACA	GAA	GTA	GT	A C	CA C	TA	bung-1. ged
917	K	L	L	R		G	T	ĸ										
	AAG								A	L	T	E	V	ν	•	₽	L	pHDMHgpm2.seq

Fig. 9E

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Alignment Report of Codon Optimization (pol).MEG, using Clustal method with PAM250 residue weight table.

	3	440										470				
3437	<u> </u>	E	E	A	E	L	E	L	A	2	N	R	E	I		NL4-3 genbank.SEQ
3437	ACA														CTA	
3435	T	E	E	A	E	L	E	L	A	E	N	R	E	I	L	pNL4-3.seq_
	ACA T	GAA E	. Gaa E	. GCA A	GAG E	CTA L	GAA E	L	GCA A	E	N N	AGG R	E	I	CTA L	pHDMHgpm2.seq
3962	ACC															pilmaigpinz.seq
,,,,																_
						3	500							_		
3482	K	E	P	٧	н	G	٧	Y	Y	D	P	5	K	D	L	NL4-3 genbank.SE
3482					CAT											-107 4 2
3480	K	E	5 5	V	H CAT	G	V	Y	Y	D	CC.9	S TCA	K	CAC D		pNL4-3.seq
3480 4007	K	E	P	V	H	G	A O	Y	Y	D	P	S	K	D	L	pHDMHgpm2.seq
	AAG													GAC		
		-										560				
		530														
3527 3527	I	A	E	I	Q CAG	X	CJC.	G	CDD.	G	Q	W TGG	T	Y Tat	-	NL4-3 genbank.SE
3525	I	A	E	I	Q	K	Q	G	Q	G	Q	W	T	Y	Q	pNL4-3.seq
	ATA											TGG	ACA	TAT		
1052	I	A	E	1	Q	K	Q	Ģ	Q	G	Q	W	T	Y	Q	pHDMHgpm2.seq
4052	ATC	GCC	GAG	ATC	CAG	AAG	CAG	GGC	CAG	GGC	CAG	TGG	ACC	TAC	CAG	
						3	590									
3572		Y	Q	E	P	F	K	· N	L	К	T	G	К	Y	A	NL4-3 genbank.SEQ
3572	ATT	TAT	_	GAG	CCA	TTT	AAA	aat	CTG	AAA	ACA	GGA	AAA	TAT	GCA	
3570	I	Y	Q	B	P	F	K	N	L	K	T	G	K	Y	A.	pNL4-3.seq
3570					CCA											-1171GI 2
1097	I	Y	Q	E	CCC	F	K	N	L	K nac	T	G	X AAA	TAC	A	pHDMHgpm2.seq
1097	ATC	TAC	CAG	GAG	ccc	110	AAG	AAC	C.5	AAG	Acc	330	~~	1,70		
	3	620									3	650				
3617	R	М	ĸ	G	A	н	T	N	D	٧	ĸ	Q	L	T	E	NL4-3 genbank.SEQ
3617	AGA	ATG													GAG	
3615	R	M	K	G	A	H	T	N	D	V	K	Q	L	T	E	pNL4-3.seq
3615			AAG K	GGT	GCC A	CAC E	ACT	AAT N	GAT D	GTG V	AAA K	Q	L	T	E	pHDMHgpm2.seq
1142	R	M ATG														P
						3	680									
662	A	V	Q	К	I	A	T	E	S	I	v	I	W	G		NL4-3 genbank.SEQ
662					ATA									GGA G		-NT 4-2 and
3660	A	v	C A A	K	I ATA	A	T	E	S AGC	I ATA	CTA V	I ATA	W TGG	-	K	pNL4-3.seq
3660 1187	GCA A	GTA V	Q	K	I	A	T	E	S	I	V	I	W	G	K	pHDMHqpm2.seq .
187	ecc .	GTG	CAG	AAG	ATC	GCC										
				•			-									

Alignment Report of Codon Optimization (pol).MEG, using Clustal method with PAM250 residue weight table.

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		710									3	740				
3707	T	P	K	F	,K	L	P	I	Q	ĸ	E	T	W	E	A	NL4-3 genbank.SE
3707	ACT	CCT	AAA												GCA	
3705	7	P	ĸ	F	K	L	P	I	Q	K	E	T	W	E.	A	pNL4-3.seq
	ACT															_
232	T	Ρ.	K	E	K	L	P	I	Q	K	E	Ţ	W	E	A	pHDMHgpm2.seq
232	ACT	ccc	AAG	TTC	AAG	CTG	CCC	ATC	CAG	AAG	GAG	ACC	TGG	GAG	GCC	
	_					3	770									•
752	· N	W	T	E	Y	W	Q	A	Ť	W	I	P	E	W	E	NL4-3 genbank.SE
752			_			TGG	-		ACC	TGG	ATT	CCT	GAG	TGG	GAG	,
750	W	W	T	E	Y	W	Q	A	T	W	1	P	E	W	E	pNL4-3.seq
750	TGG	TGG	ACA	GAG	TAT	TGG	CAA	GCC	ACC	TGG	ATT	CCT	GAG	TGG	GAG	
277	W	W	T	E	Y	W	Q	A	T	W	I	P	E	W	E	pHDMHgpm2.seq
277	TGG	TGG	ACC	GAG	TAC	TGG	CAG	GCC	ACC	TGG	ATC	CCC	GAG	TGC	GAG	
		800										830		_		
										_	78	٠		_		
797	F	v	N	T	5	E E	L	V -	K	L		The	e co	L	E	NL4-3 genbank.SE
	TTT					P		V	K	L	W	Y	Q	L	E E	-177.4.3
795	F TTT	V	N	T	P	-	L						_			pNL4-3.seq
195 322	F	V	. N	ACC	P	P	L	A	· K	L	W	Y	δ.	L	E	pHDMHqpm2.seq
	TTC															phorngpmz.seq
							860				Y	v				MT 4 2 manhanis CE
842	K	3	P.	I	I	G	A	E	T	F TTC	-		D CAT	3	A	NL4-3 genbank.SE
842			2	I	AIA I	GGA G	A	E	T	F	, X	V	D	G	A	pNL4-3.seq
840	K AAA	E		_							_		-		GCA	pana-3.seq
367	K	E	P	I	I	G	A	E	T	F	Y	v	5	G	A	pHDMHgpm2.sec
	AAG							_	_							5.101219pmm 1004
												•				
	3	890									3	920				
887	A	N	R	E	T	ĸ	L	G	К	A	G	Y	V	T	D	NL4-3 genbank.SE
887	GCC					AAA									GAC	
885	A	N	R	E	T	K	L	G	K	A	G	Y	v	T	D	pNL4-3.seq
885						AAA										
412	A	N	R	E	T	K	<u>.</u>	G	K	A	G	Y	V	T	D	pHDMHgpm2.seq
12	GCC	AAC	CGC	GAG	ACC	AAG	CTG	GGC	AAG	نابان	GGC	IAC	GIG	ACC	UAC	
	—					3:	950									
932	R	G	R	Q	ж	v		P	5	T	D	T	ī	N	0	NL4-3 genbank.520
932	AGA			-		GTT		_		ACG	GAC		ACA		_	
30	R	G	R	Q	K	٧	٧	P	L	T	Đ	T	T	N	Q	pNL4-3.seq
30	AGA					GTT	GTC	CCC	CTA	ACG	GAC	ACA	ACA	AAT	CAG	· ·
157	R	G	R	Q	K	v	V	P	L	T	D	T	T	N	Q	pHDMHgpm2.seq
57	cec	GGC	CGC	CAG	AAG	GTG	GTG	ccc	CTG	ACC	GAC	ACC	ACC	AAC	CAG	

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Alignment Report of Codan Optimization (pol).MEG, using Clustal method with PAM250 residue weight table.

		3980										1010				
3977	К	T	E	L	Q	A	I	Ħ	L	A	L	Q	D	5	G	
3977		ACT		TTA											GGA	
3975	K	T	E	L	Q	A	I	H	L	A	L	Q CAG	D CDT	S	G	pNL4-3.seq
3975 4502		ACT T	GAG E	TTA L	CAA Q	A GCA	. ATI	H	L	A	L	Q	. פעז	S	G G	pHDMHgpm2.seq
	K Aag															
1502		7100														- -
						4	040									
4022	L	E	v	N	1	٧	T	D	. s	Q	Y	A	L	G	I	NL4-3 genbank.SEC
4022	TTA			AAC												
4020	L	E	٧	N	I	V	T	D	S	Q	Y	A	L	G	I	pNL4-3.seq
4020		GAA E	GTA V	AAC N	ATA	. GTG	ACA T	. GAU	S	Q	Y	A	. IIG	G	I	pHDMHgpm2.seq
4547	L CTG															bunundhur. sed
		-										т-				-
		070										100				•
4067	1	Q	A	Q	P	D	X	S	E	S	2	L	V	S	Q	NL4-3 genbank.SEQ
4067	ATT		. GCA A	CAA	CCA P	GAT D	AAG	AGT	E	S	E	L	A CLC	AGI S	O	pNL4-3.seg
4065		Q AA		Q CAA												prot-3.3eq
4592	ï	Q	A.	Q	P	D	K	S	E	S	E	L	v	S	Q	pHDMHgpm2.seq
4592	ATC	CAG	GCC	CAG	CCC	GAC	AAG	TCC	GAG	TCC	GAG	CTG	GTG	TCC	CAG	
							130									
4112		I	E	Q	L	I	R	ĸ	E	K	v	Y	L	A	W	NL4-3 genbank.SEQ
4112																not o genounitons
4110	I	I	E	Q	L	1	K	K	E	ĸ	ν	Y	L	A	W	pNL4-3.seq
4110	ATA	ATA	GAG	CAG	TTA	ATA	AAA	AAG	Gλλ			TAC		GCA		
4637	I	I	Ē	Q	L	I	ĸ	K	Ē	K	٧	Y	L	Α	W	pHDMHgpm2.seq
4637	ATC	ATC	GAG	CAG	CTG	ATC	AAG	AAG	GAG	AAG	GTG	TAC	CTG	GCC	TGG	
	4	160									4	190				
4157		P	A	н	К	<u> </u>	I	G	G	N	E	-	v	D	G	NL4-3 genbank.SEQ
4157				CAC							_					
4155	٧	P	A	H	K	G	I	G	G	N	E	Q	٧	D	K	pNL4-3.seq
4155	GTA	CCA	GCA	CAC	AAA	GGA										
4682	V	P	A	H	K	G	I	G	G	N	E	Q	V	D	K	pHDMHgpm2.seq
4682	GTG	CCC	GCC	CAC	AAG	GGC	ATC	GGC	GGC	AAC	GAG	CAG	GTG	GAC	AAG	
						4	220									
4202	-L	v	3		G	ī	R	к		L	F	L	D	G	ī	NL4-3 genbank.SEQ
4202	TTG	-	-		GGA	_		-		CTA	_			GGA	_	
4200	L	v	S	A	Ğ.	I	R	ĸ	V	L	P	L	D	G	I	pNL4-3.seq
4200	TTG	GTC	AGT	GCT	GGA	ATC	AGG	AAA	GTA					GGA		
4727	L	v	S	A	G	I	R	K	V	L	F	L	D	G	I	pHDMHgpm2.seq
4727	CTG	GTG	TCC	GCC	GGC	ATC	CCC	AAG	GTG	CTG	TTC	CTG	GAC	GGC	ATC	

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Alignment Report of Codon Optimization (pol).MEG, using Clustal method with PAM250 residue weight table.

4250 47 D K A Q E E H E K Y H S N W R 77 GAT ANG GCC CAA GAA CAA CAT CAT GAG AAA TAT CAC AGT AAT TGG AGA 45 D K A Q E E H E K Y H S N W R 45 D K A Q E E H E K Y H S N W R 45 GAT ANG GCC CAA GAA GAA CAT GAG AAA TAT CAC AGT AAT TGG AGA 45 D K A Q E E H E K Y H S N W R 45 GAT ANG GCC CAA GAA GAA CAT GAG AAA TAT CAC AGT AAT TGG AGA 45 GAT ANG GCC CAA GAG GAG CAC GAG AAA TAT CAC AGT AAT TGG AGA 72 D K A Q E E H E K Y H S N W R 72 D K A Q E E H E K Y H S N W R 74 D W W W W W W W W W W W W W W W W W W		,															
47 GAT AAG SCC CAA GAA GAA CAT GAG AAA TAT CAC AGT AAT TGG AGA 45 D K A Q E E H E K Y H S N W R 45 GAT AAG GCC CAA GAA GAA CAT GAG AAA TAT CAC AGT AAT TGG AGA 72 D K A Q E E H E K Y H S N W R 72 GAC AAG GCC CAA GAA GAA CAT GAG AAA TAT CAC AGT AAT TGG AGA 73 D K A Q E E H E K Y H S N W R 74 PHDMHgpm2.sec 4310 4310 4310 4310 A M A S D F N L P P V V A K E 9 PHDMHgpm2.sec 60 A ATG GCT AGT GAT TTT AAC CTA CCA CCT GTA GTA GCA AAA GAA 90 A M A S D F N L P P V V A K E 90 GCA ATG GCT AGT GAT TTT AAC CTA CCA CCT GTA GTA GCA AAA GAA 17 A M A S D F N L P P V V A K E 17 AC CAT GCT AGT GAT TTT AAC CTA CCA CCT GTA GTA GCA AAA GAA 17 A M A S D F N L P P V V A K E 17 AC CAT GCT AGT GAT TTT AAC CTA CCA CCT GTA GTA GCA AAA GAA 18 A A S D F N L P P V V A K E 18 PHDMHgpm2.sec 4340 4370 50 K C Q L K G E A M 50 M M A S C D K C Q L K G E A M 50 M M A S C D K C Q L K G E A M 50 M M A S C D K C Q L K G E A M 50 M M A S C D K C Q L K G E A M 50 M M A S C D K C Q L K G E A M 50 M M A S C D K C Q L K G E A M 50 M M A S C D K C Q L K G E A M 50 M M A S C D K C Q L K G E A M 50 M M A S C D K C Q L K G E A M 50 M M M M M M M M M M M M M M M M M M M			1250														
45	4247	D	K	A	Q	E	Е	H	E	К	Y	Н	s	N	M	R	NL4-3 genbank.
4310 92	4247	GAT	AAG	GCC	CAA	A GAA	GAA	CA1	GAG	AA.	TAT	CAC	AG?	: AA1	TGO) AGA	• 1
72 D K A Q E E H E K Y H S N W R PHDMHgpm2.seq 4310 92 AA M A S D F N L P P V V A K E PHL4-3.seq 90 GCA ATG GCT AGT GAT TIT AAC CTA CCA CCT GTA GTA GCA AAA GAA 17 A M A S D F N L P P V V A K E PHDMHgpm2.seq 18 GCA ATG GCT AGT GAT TIT AAC CTA CCA CCT GTA GTA GCA AAA GAA 17 A M A S D F N L P P V V A K E PHDMHgpm2.seq 18 GCA ATG GCT AGT GAT TIT AAC CTA CCA CCT GTA GTA GCA AAA GAA 17 A M A S D F N L P P V V A K E PHDMHgpm2.seq 18 GCA ATG GCT AGT GAT TIT AAC CTA CCA CCT GTA GTA GCA AAA GAA 18 A M A S D F N L P P V V A K E PHDMHgpm2.seq 19 GCA ATG GCC TCC GAC TTC AAC CTG CCC CCC GTG GTG GCC AAG GAG 4340 4370 37 ATA GTA GCC AGC TGT GAT AAA TGT CAG CTA AAA GGG GAA GCC ATG 38 ATA GTA GCC AGC TGT GAT AAA TGT CAG CTA AAA GGG GAA GCC ATG 39 I V A S C D K C Q L K G E A M PAL4-3.seq 30 ATA GTA GCC AGC TGT GAT AAA TGT CAG CTA AAA GGG GAA GCC ATG 30 ATA GTA GCC AGC TGT GAT AAA TGT CAG CTA AAA GGG GAA GCC ATG 31 I V A S C D K C Q L K G E A M PAL4-3.seq 32 ATC GTG GCC TCC TGC GAC AAG TGC CAG CTG AAG GGC GAG GCC ATG 32 I V A S C D K C Q L K G E A M PAL4-3.seq 33 ATA GTA GCC AGC TGT GAT AAA TGT CAG CTA AAA GGG GAA GCC ATG 4400 82 H G Q V D C S P G I W Q L D C PAL4-3.seq 90 CAT GGA CAA GTA GAC TGT AGC CCA GGA ATA TGG CAG CTA GAT TGT 07 H G Q V D C S P G I W Q L D C PAL4-3.seq 17 CAC GGC CAG GTG GAC TGC CCC GGC ATC TGG CAG CTA GAT TGT 18 GC CAT GGA CAA GTA GAC TGT AGC CCC GGC ATC TGG CAG CTA GAT TGT 19 CAC GGC CAG GTG GAC TGC CCC CCG GGC ATC TGG CAG CTA GAT TGT 10 T H L E G K V I L V A V H V A NL4-3 genbank 12 ACA CAT TTA GAA GGA AAA GTT ATC TTG GTA GCA GTT CAT GTA GCC 14400 4460 27 T H L E G K V I L V A V H V A NL4-3 genbank 18 C CAT GGA CAA GTA GGA AAA GTT ATC TTG GTA GCA GTT CAT GTA GCC 19 CAC GGA CAA GTA AGG CAAA GTT ATC TTG GTA GCA GTT CAT GTA GCC 19 CAC GGA CAA GTA AGA GTA ATT CCA GCA GAA ACA GGG CAA 19 C ATT GGA TAT ATA GAA GAA GTA ATT CCA GCA GAA ACA GGG CAA 19 C ATT GGA TAT ATA GAA GAA GTA ATT CCA GCA GAA ACA GGG CAA 19 C ATT GGA TATA ATA GAA GAA GTA ATT CCA GCA GAA GAA GAA GGC CAA 19 C ATT GGA TATA ATA GAA GCA GAA	4245				-	_	_									R	pNL4-3.seq
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4310 92 A M A S D F N L P P V V A K E NL4-3 genbank 92 GCA ATG GCT AGT GAT TTT AAC CTA CCA CCT GTA GTA GCA AAA GAA 90 A M A S D F N L P P V V A K E PL4-3.seq 90 GCA ATG GCT AGT GAT TTT AAC CTA CCA CCT GTA GCA AAA GAA 17 A M A S D F N L P P V V A K E PL4-3.seq 91 GCA ATG GCT AGT GAT TTT AAC CTA CCA CCT GTA GCA AAA GAA 17 A M A S D F N L P P V V A K E PL4-3.seq 91 GCA ATG GCC TCC GAC TTC AAC CTG CCC CCC GTG GTG GCC AAG GAG 4340 4370 4370 11 V A S C D K C Q L K G E A M NL4-3 genbank 935 ATA GTA GCC AGC TGT GAT AAA TGT CAG CTA AAA GGG GAA GCC ATG 935 I V A S C D K C Q L K G E A M NL4-3.seq 94400 AND A S C D K C Q L K G E A M NL4-3 genbank 980 CAT GGA CAA GTA GAC TGT AGC CCA GGA ATA TGG CAG CTA GAT TGT 980 CAT GGA CAA GTA GAC TGT AGC CCA GGA ATA TGG CAG CTA GAT TGT 980 CAT GGA CAA GTA GAC TGT AGC CCA GGA ATA TGG CAG CTA GAT TGT 980 CAT GGA CAA GTA GAC TGT AGC CCA GGA ATA TGG CAG CTA GAT TGT 980 CAT GGA CAA GTA GAC TGT AGC CCA GGA ATA TGG CAG CTA GAT TGT 980 CAT GGA CAA GTA GAC TGT AGC CCA GGA ATA TGG CAG CTA GAT TGT 980 CAT GGA CAA GTA GAC TGT AGC CCA GGA ATA TGG CAG CTA GAT TGT 980 CAT GGA CAA GTA GAC TGT AGC CCA GGA ATA TGG CAG CTA GAT TGT 980 CAT GGA CAA GTA GAC TGT AGC CCA GGA ATA TGG CAG CTA GAT TGT 980 CAT GGA CAA GTA GAC TGT AGC CCA GGA ATA TGG CAG CTA GAT CTC 980 CAT GGA CAA GTA GAC TGT AGC CCA GGA ATA TGG CAG CTA GAT CTC 980 CAT GGA CAA GTA GAC TGT AGC CCA GGA CAG CTA GTA GCC 980 CAT GGA CAA GTA GAC AGA GTA ATT CTG GTA GCC GTG CAC GTG GCC 980 CAG GGA GGA GGA GGA GTA ATC CTG GTA GCC GTG CAC GTG GCC 981 T H L E G K V I L V A V H V	772				-	_	_		_								pHDMHgpm2.seq
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12 S G Y I E A E V I P A E T G Q NL4-3 genbank. 12 AGT GGA TAT ATA GAA GCA GAA GTA ATT CCA GCA GAG ACA GGG CAA 10 S G Y I E A E V I P A E T G Q pNL4-3.seq 10 AGT GGA TAT ATA GAA GCA GAA GTA ATT CCA GCA GAG ACA GGG CAA 17 S G Y I E A E V I P A E T G Q pHDMHgpm2.seq	952	ACC	CAC	CTG	GAG	GGC	AAG	GTG	ATC	CTG	GTG	GCC	GTG	CAC	GTG	GCC	
12 S G Y I E A E V I P A E T G Q NL4-3 genbank. 12 AGT GGA TAT ATA GAA GCA GAA GTA ATT CCA GCA GAG ACA GGG CAA 10 S G Y I E A E V I P A E T G Q pNL4-3.seq 10 AGT GGA TAT ATA GAA GCA GAA GTA ATT CCA GCA GAG ACA GGG CAA 17 S G Y I E A E V I P A E T G Q pHDMHgpm2.seq																	
22 AGT GGA TAT ATA GAA GCA GAA GTA ATT CCA GCA GAG ACA GGG CAA 10 S G Y I E A E V I P A E T G Q pNL4-3.seq 10 AGT GGA TAT ATA GAA GCA GAA GTA ATT CCA GCA GAG ACA GGG CAA 17 S G Y I E A E V I P A E T G Q pHDMHgpm2.seq																	
OSGYIE AEVIPAETG Q pNL4-3.seq OAGT GGA TAT ATA GAA GCA GAA GTA ATT CCA GCA GAG ACA GGG CAA OSG YIE AEVIPAETG Q pHDMHgpm2.seq	172	_		-	-			_	-				_			_	NL4-3 genbank.S
0 AGT GGA TAT ATA GAA GCA GAA GTA ATT CCA GCA GAG ACA GGG CAA 17 S G Y I E A E V I P A E T G Q pHDMHgpm2.seq	472																
7 SGYIEAEVIFAETGQpHDMHgpm2.seq	170		_			-			-								bur4-3.sed
	70																
77 THE GGE THE ATC GAG GEE GAG GTG ATT EET GEE GAG ACE GGE CAG	197	-	_	_	_	_		_		_							parangpmz.seq
	97	ICC	GGC	TAC	ATC	GAG	GCC	GAG	GTG	ATC	CCC	GCC	GAG	ACC	GGC	CAG	

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Alignment Report of Codon Optimization (pol).MEG, using Clustel method with PAM250 residue weight table.

4517 GAA ACA GCA TAC TTC CTC TTA AAA TTA GCA GGA AGA TGG CCA GTA 4515 E T A Y F L L K L A G R W P V PNI 4515 GAA ACA GCA TAC TTC CTC TTA AAA TTA GCA GGA AGA TGG CCA GTA 5042 E T A Y F L L K L A G R W P V PNI 5042 GAG ACC GCC TAC TTC CTG CTG AAG CTG GCC GGC CGC TGG CCC GTG 4580 4560 4560 K T V H T D N G S N F T S T T NL4 4560 K T V H T D N G S N F T S T T PNI 4560 AAA ACA GTA CAT ACA GAC AAT GGC AGC AAT TTC ACC AGT ACT ACA 4560 AAA ACA GTA CAT ACA GAC AAT GGC AGC AAT TTC ACC AGT ACT ACA 4560 AAA ACA GTA CAT ACA GAC AAT GGC AGC AAT TTC ACC AGT ACT ACA 5087 K T V H T D N G S N F T S T T PNI 4560 AAA ACA GTA CAT ACA GAC AAT GGC AGC AAT TTC ACC AGT ACT ACA 5087 K T V H T D N G S N F T S T T PNI 4610 4640 4610 4640 4610 7 V K A A C W W A G I K Q E F G NL4 4607 GTT AAG GCC GCC TGT TGG TGG GCG GGG ATC AAG CAG GAA TTT GGC	4-3 genbank.SEG .4-3.seq OMHgpm2.seq 1-3 genbank.SEG .4-3.seq
4517 E T A Y F L L K L A G R W P V NL4 4517 GAA ACA GCA TAC TTC CTC TTA AAA TTA GCA GGA AGA TGG CCA GTA 4515 E T A Y F L L K L A G R W P V PNI 4515 GAA ACA GCA TAC TTC CTC TTA AAA TTA GCA GGA AGA TGG CCA GTA 4515 GAA ACA GCA TAC TTC CTC TTA AAA TTA GCA GGA AGA TGG CCA GTA 5042 E T A Y F L L K L A G R W P V PHI 5042 GAG ACC GCC TAC TTC CTG CTG AAG CTG GCC GGC CGC TGG CCC GTG 4580 4562 R T V H T D N G S N F T S T T NL4 4562 AAA ACA GTA CAT ACA GAC AAT GGC AGC AAT TTC ACC AGT ACT ACA 4560 R T V H T D N G S N F T S T T PNI 4560 AAA ACA GTA CAT ACA GAC AAT GGC AGC AAT TTC ACC AGT ACT ACA 4560 AAA ACA GTA CAT ACA GAC AAT GGC AGC AAT TTC ACC AGT ACT ACA 4560 AAA ACA GTA CAT ACA GAC AAT GGC AGC AAT TTC ACC AGT ACT ACA 4560 AAA ACA GTA CAT ACA GAC AAT GGC AGC AAT TTC ACC AGT ACT ACA 4560 AAA ACA GTA CAT ACA GAC AAT GGC AGC AAT TTC ACC AGT ACT ACA 4610 4640 4610 4640 4607 V K A A C W W A G I K Q E F G NL4 4607 GTT AAG GCC GCC TGT TGG TGG GCG GGG ATC AAG CAG GAA TTT GCC	A-3.seq MHgpm2.seq 1-3 genbank.SEq 4-3.seq
4517 GAA ACA GCA TAC TTC CTC TTA AAA TTA GCA GGA AGA TGG CCA GTA 4515 E T A Y F L L K L A G R W P V PNI 4515 GAA ACA GCA TAC TTC CTC TTA AAA TTA GCA GGA AGA TGG CCA GTA 5042 E T A Y F L L K L A G R W P V PNI 5042 GAG ACC GCC TAC TTC CTG CTG AAG CTG GCC GGC CGC TGG CCC GTG 4580 4560 K T V H T D N G S N F T S T T NL4 4562 AAA ACA GTA CAT ACA GAC AAT GGC AGC AAT TTC ACC AGT ACT ACA 4560 K T V H T D N G S N F T S T T PNI 4560 AAA ACA GTA CAT ACA GAC AAT GGC AGC AAT TTC ACC AGT ACT ACA 4560 AAA ACA GTA CAT ACA GAC AAT GGC AGC AAT TTC ACC AGT ACT ACA 5087 K T V H T D N G S N F T S T T PNI 4560 AAA ACA GTA CAT ACA GAC AAT GGC AGC AAT TTC ACC AGT ACT ACA 5087 K T V H T D N G S N F T S T T PNI 4607 AAG ACC GTG CAC ACC GAC AAC GGC TCC AAC TTC ACC TCC ACC 4610 4640 4607 V K A A C W W A G I K Q E F G NL4 4607 GTT AAG GCC GCC TGT TGG TGG GCG GGG ATC AAG CAG GAA TTT GCC	A-3.seq MHgpm2.seq 1-3 genbank.SEq 4-3.seq
4515 E T A Y F L L K L A G R W P V PNI 4515 GAA ACA GCA TAC TTC CTC TTA AAA TTA GCA GGA AGA TGG CCA GTA 5042 E T A Y F L L K L A G R W P V PHI 5042 GAG ACC GCC TAC TTC CTG CTG AAG CTG GCC GGC CGC TGG CCC GTG 4580 4580 4580 4580 A580 A680 A780 A	OMHgpm2.seq 1-3 genbank.SE(
4515 GAA ACA GCA TAC TTC CTC TTA AAA TTA GCA GGA AGA TGG CCA GTA 5042 E T A Y F L L K L A G R W P V MT P L L K L A G R W P V MT P L L K L A G R W P V MT P L L K L A G R W P V MT P L L K L A G R W P V MT P L L K L A G R W P V MT P L L K L A G R W P V MT P L L K L A G R W P V MT P L L K L A G R W P V MT P L L K L A G R W P V MT P L L K L A G R W P V MT P L L K L A G R W P V MT P L L K L A G R W P V MT P L L K L A G R W P V MT P L L K L A G R W P V MT P L L K L A G R W P V MT P L L K L A G R R W P V MT P L L K L A G R R W P V MT P L L K L A G R W P V MT P L L L K L A G R W P V MT P L L K L A G R W P V MT P L L K L A G R W P V MT P L L K L A G R W P V MT P L L K L A G R C G G G G G G G G G G G G G G G G G	1-3 genbank.SE(4-3.seq
4580 4580 4580 4582 R T V H T D N G S N F T S T T NL4 4562 AAA ACA GTA CAT ACA GAC AAT GGC AGC AAT TTC ACC AGT ACT ACA 4560 R T V H T D N G S N F T S T T PNI 4560 AAA ACA GTA CAT ACA GAC AAT GGC AGC AAT TTC ACC AGT ACT ACA 4560 AAA ACA GTA CAT ACA GAC AAT GGC AGC AAT TTC ACC AGT ACT ACA 5087 AT V H T D N G S N F T S T T PHI 5087 AAG ACC GTG CAC ACC GAC AAC GGC TCC AAC TTC ACC TCC ACC 4610 4610 4640 4640 V K A A C W W A G I K Q E F G NL4 4607 GTT AAG GCC GCC TGT TGG TGG GGG GGG ATC AAG CAG GAA TTT GGC	1-3 genbank.SE(4-3.seq
4580 4562 R T V H T D N G S N F T S T T NL4 4562 AAA ACA GTA CAT ACA GAC AAT GGC AGC AAT TTC ACC AGT ACT ACA 4560 R T V H T D N G S N F T S T T PNI 4560 AAA ACA GTA CAT ACA GAC AAT GGC AGC AAT TTC ACC AGT ACT ACA 5087 R T V H T D N G S N F T S T T PHI 5087 AAG ACC GTG CAC ACC GAC AAC GGC TCC AAC TTC ACC TCC ACC 4610 4610 4640 4640 V K A A C W W A G I K Q E F G NL4 4607 GTT AAG GCC GCC TGT TGG TGG GCG GGG ATC AAG CAG GAA TTT GCC	4-3.seq
4562	4-3.seq
4562 AAA ACA GTA CAT ACA GAC AAT GGC AGC AAT TTC ACC AGT ACT ACA 4560 R T V H T D N G S N F T S T T PNI 4560 AAA ACA GTA CAT ACA GAC AAT GGC AGC AAT TTC ACC AGT ACT ACA 5087 R T V H T D N G S N F T S T T PHI 5087 AAG ACC GTG CAC ACC GAC AAC GGC TCC AAC TTC ACC TCC ACC 4610 4640 4607 V K A A C W W A G I K Q E F G NL4 4607 GTT AAG GCC GCC TGT TGG TGG GGG GGG ATC AAG CAG GAA TTT GGC	4-3.seq
4560 K T V H T D N G S N F T S T T PNI 4560 AAA ACA GTA CAT ACA GAC AAT GGC AGC AAT TTC ACC ATT ACA 5087 K T V H T D N G S N F T S T T T PHE 5087 AAG ACC GTG CAC ACC GAC AAC GGC TCC AAC TTC ACC TCC ACC 4610 4640 V K A A C W W A G I K Q E F G NL4 4607 GTT AAG GCC GCC TGT TGG TGG GCG GGG ATC AAG CAG GAA TTT GCC	
4610 4610 4610 4610 4610 4610 4640 4607 V K A A C W W A G I K Q E F G NL4 4607 GTT AAG GCC GCC TGT TGG TGG GGG GGG ATC AAG CAG GAA TTT GCC	
4610 4610 4610 V K A A C W W A G I K Q E F G NL4 4607 CTT AAG GCC GCC TGT TGG GGG GGG ATC AAG CAG GAA TTT GCC	MHgpm2.seq
AAG ACC GTG CAC ACC GAC AAC GGC TCC AAC TTC ACC TCC ACC ACC 4610 4640 V K A A C W W A G I K Q E F G NL4 4607 GTT AAG GCC GCC TGT TGG TGG GCG GGG ATC AAG CAG GAA TTT GGC	mngpmz.seq
4610 4640 4607 V K A A C W W A G I K Q E F G NL4 4607 GTT AAG GCC GCC TGT TGG TGG GCG GGG ATC AAG CAG GAA TTT GGC	
4610 4640 4607 V K A A C W W A G I K Q E F G NL4 4607 GTT AAG GCC GCC TGT TGG TGG GGG ATC AAG CAG GAA TTT GGC	
4607 GTT AAG GCC GCC TGT TGG TGG GCG GGG ATC AAG CAG GAA TTT GGC	
	l-3 genbank.SE
ACOS V K A A C W W A G T K O E F G PNT	
	.4-3.seq
4605 GTT AAG GCC GCC TGT TGG TGG GCG GGG ATC AAG CAG GAA TTT GGC	wer
	MHgpm2.seq
5132 GTG AAG GCC GCC TGC TGG TGG GCC GGC ATC AAG CAG GAG TTC GGC	
4670	
7004 2 4 4 " " " " " " " " " " " " " " " "	I-3 genbank.SEG
4652 ATT CCC TAC AAT CCC CAA AGT CAA GGA GTA ATA GAA TCT ATG AAT	
	14-3.seg
4650 ATT CCC TAC AAT CCC CAA AGT CAA GGA GTA ATA GAA TCT ATG AAT	Milmon? and
	MHgpm2.seq
5177 ATC CCC TAC AAC CCC CAG TCC CAG GGC GTG ATC GAG TCC ATG AAC	
4700 4730	
	l-3 genbank.SEG
4697 AAA GAA TTA AAG AAA ATT ATA GGA CAG GTA AGA GAT CAG GCT GAA	
	.4-3.seq
4695 AAA GAA TTA AAG AAA ATT ATA GGA CAG GTA AGA GAT CAG GCT GAA	Wilson? east
	MHgpm2.seq
5222 AAG GAG CTG AAG AAG ATC ATC GGC CAA GTC CGC GAC CAG GCC GAG	
4760	
	-3 genbank.SEQ
4742 CAT CTT AAG ACA GCA GTA CAA ATG GCA GTA TTC ATC CAC AAT TTT	
	.4-3. seq
4740 CAT CTT AAG ACA GCA GTA CAA ATG GCA GTA TTC ATC CAC AAT TTT	na:
	MHgpm2.seq
5267 CAC CTG AAG ACC GCC GTG CAG ATG GCC GTG TTC ATC CAC AAC TTC	

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Alignment Report of Codon Optimization (pol).MEG, using Clustal method with PAM250 residue weight table.

																, worght about.
•		479	0									4820				
4787					G	I	G	G	Y	s	A	G	Z			
4787					G GGG								G GA	A AG	A AT	A
4785		-			G		G	G	Y	S	A	G	E	R		pNL4-3.seq
4785 5312					e eee											
		-			G GGG	I	. cc	G	Y The	S	A	·G	E	R		pHDMHgpm2.seq
3312		3 C3	C 7/1		. 666	, MIC	. 66	- 664	_ IA	. 100	. 600	. 666	- GA	3 (G	CAT	•
							4850									-
4832		D	I	I	A	T	D	1	Q	T	ĸ	Е	L	Q	К	NL4-3 genbank.SE
4832					A GCA						AA	A GA	TT?	A CA	AA A	١
4830		D	I.	_ I	Α	T	D	I	Q	T	K	E	L	Q	K	pNL4-3.seq
4830					A GCA											1
5357 5357		D עבו	I TAT	I Otac	A GCC	T	C D	I Tro	Q Cac	T	K	E .	L	Q Q	K	pHDMHgpm2.seq
							- GAG		· Grid		7/10	o Grac	, ст	. CA	J AAL	•
		4880				_						4910				_
4877	Q	1	T	X	I	Q	И	7	R	V	Y	Y	R	D	S	
4877					L ATT								AGG	GAG	AGC	
4875	Q	I	T	ĸ	I	Q	N	F	R	V	Y	Y	R	D	S	pNL4-3.seq
4875					ATT										AGC	
5402 5402	_	I TATE	T ACC	K Tabo	I ATC	C)C	N	E	R	V	Y	Y	R	D		pHDMHgpm2.seq
J402	CAG	, ,,,,,		- 200	AIC	CAG	AAC	110	CGC	GIG	TAC	TAC	CGG	GAC	TCC	•
						4	940				,					-
922	R	D	P	v	W	К	G	5	A	К	L	L	W	к	G	NL4-3 genbank.SEQ
1922		GA:		GTT	TGG			CCA	GCA	AAG	CTC	CTC	TGG	AAA	GGT	
1920	R	D	P	٧	W	K	G	Р	A	K	L	L	W	K	G	pNL4-3.seq
1920					TGG											
5447 5447	R	D	P	. ~~~	W	X	G	P	A	K	L	L	W	К	G	pHDMHgpm2.seq
7447	CGC	GAC		GIG	TGG	AAG	GGC	CCC	GCC	AAG	CTG	CTG	TGG	AAG	GGC	
	4	1970									5	000				•
967	E	G	A	v	v	I	Q	D	N	S	D	I	К	٧	٧	NL4-3 genbank.SEQ
967		GGG			GTA											
965	E	G	A	V	٧	I	Q	D	N	5	D	I	K	v	٧	pNL4-3.seq
965 492	GAA	GGG	GCA A	. GTA	GTA V											
492					GTG	I ATC	CAG.	באר. D	N	S TCC	EDC.	I	K	V GTG	A.	pHDMHgpm2.seq
				0.0				unc	M	100	unc	A10	~~~	010	316	
						50	30									
012	P	Ŗ	R	K	A	ĸ	I	1	R	D	Y	G	ĸ	Q.	М	NL4-3 genbank.SEQ
012	CCA	AGA	AGA	AAA	GCA	AAG .	ATC	ATC	AGG	GAT	TAT	GGA	AAA	CAG	ATG	,
010	P	R	R	K	A	K	I	Į	R	D	Y	G	ĸ	Q	M	pNL4-3.seq
010					GCA									CAG	ATG	
537	P	R	R	K	A	K	I	I	R	D	Y	G	ĸ	Q	M	pHDMHgpm2.seq
537	ccc	CGC	CGC	AAG	GCC .	AAG .	ATC	ATC	CGC	GAC	TAC	GGC	AAG	CAG	ATG	

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Alignment Report of Codon Optimization (pol), MEG, using Clustal method with PAM250 residue weight table.

		060									5	090			
5057	A	G	D		С		А	s	R	Q	D	E	D	τ	NL4-3 genbank.SEQ
5057	GCA	GGT	gat	GAT	tgt	GTG	GCA	AGT	AGA	CAG	gat	GAG	GAT	TAA.	_
5055	A	G	Ð	D	C	V	A		R		D	E	D		pNL4-3.seq
5055	GCA	GGT	GAT	GAT	TGT	GTG	GCA	AGT	AGA	CAG	GAT	GAG	GAT	TAK	-
5582	Α	G	D	D	C	٧	A	S	R	Q	D	E	D	200	pHDMHqpm2.seq
5582	GCC	GGC	GAC	GAC	TGC	GTG	GCC	TCC	CGC	CAG	CAC	GAG	GAC	TAA	

Fig. 9L

AGCTTGGCCC	ATTGCATACG	TTGTATCCAT	ATCATAATAT	GTACATTTAT	ATTGGCTCAT	60
GTCCAACATT	ACCGCCATGT	TGACATTGAT	TATTGACTAG	TTATTAATAG	TAATCAATTA	120
CGGGGTCATT	AGTTCATAGC	CCATATATGG	AGTTCCGCGT	TACATAACTT	ACGGTAAATG	180
GCCCGCCTGG	CTGACCGCCC	AACGACCCCC	GCCCATTGAC	GTCAATAATG	ACGTATGTTC	240
CCATAGTAAC	GCCAATAGGG	ACTTTCCATT	GACGTCAATG	GGTGGAGTAT	TTACGGTAAA	300
CTGCCCACTT	GGCAGTACAT	CAAGTGTATC	ATATGCCAAG	TACGCCCCCT	ATTGACGTCA	360
ATGACGGTAA	ATGGCCCGCC	TGGCATTATG	CCCAGTACAT	GACCTTATGG	GACTTTCCTA	420
CTTGGCAGTA	CATCTACGTA	TTAGTCATCG	CTATTACCAT	GGTGATGCGG	TTTTGGCAGT	480
ACATCAATGG	GCGTGGATAG	CGGTTTGACT	CACGGGGATT	TCCAAGTCTC	CACCCCATTG	540
ACGTCAATGG	GAGTTTGTTT	TGGCACCAAA	ATCAACGGGA	CTTTCCAAAA	TGTCGTAACA	600
ACTCCGCCCC	ATTGACGCAA	ATGGGCGGTA	GGCGTGTACG	GTGGGAGGTC	TATATAAGCA	660
GAGCTCGTTT	AGTGAACCGT	CAGATCGCCT	GGAGACGCCA	TCCACGCTGT	TTTGACCTCC	720
ATAGAAGACA	CCGGGACCGA	TCCAGCCTCC	CCTCGAAGCT	GATCCTGAGA	ACTTCAGGGT	780
GAGTCTATGG	GACCCTTGAT	GTTTTCTTTC	CCCTTCTTTT	CTATGGTTAA	GTTCATGTCA	840
TAGGAAGGGG	AGAAGTAACA	GGGTACACAT	ATTGACCAAA	TCAGGGTAAT	TTTGCATTTG	900
TAATTTTAAA	AAATGCTTTC	TTCTTTTAAT	ATACTTTTTT	GTTTATCTTA	TTTCTAATAC	960
TTTCCCTAAT	CICITICITI	CAGGGCAATA	ATGATACAAT	GTATCATGCC	TCTTTGCACC	1020
ATTCTAAAGA	ATAACAGTGA	TAATTTCTGG	GTTAAGGCAA	TAGCAATATT	TCTGCATATA	1080
AATATTTCTG	CATATAAATT	GTAACTGATG	TAAGAGGTTT	CATATTGCTA	ATAGCAGCTA	1140
CAATCCAGCT	ACCATTCTGC	TTTTATTTTA	TGGTTGGGAT	AAGGCTGGAT	TATTCTGAGT	1200
CCAAGCTAGG	CCCTTTTGCT	AATCATGTTC	ATACCTCTTA	TCTTCCTCCC	ACAGCTCCTG	1260
GGCAACGTGC	TGGTCTGTGT	GCTGGCCCAT	CACTTTGGCA	AAGAATTCTA	GACTGCCATG	1320
GCCCCCCCC	CCTCCGTGCT	GTCCGGCGGC	GAGCTGGACA	AGTGGGAGAA	GATCCGCCTG	1380
CGCCCCGGCG	GCAAGAAGCA	GTACAAGCTG	AAGCACATCG	TGTGGGCCTC	CCGCGAGCTG	1440
GAGCGCTTCG	CCGTGAACCC	CGGCCTGCTG	GAGACCTCCG	AGGGCTGCCG	CCAGATCCTG	1500
GGCCAGCTGC	AGCCCTCCCT	GCAAACCGGC	TCCGAGGAGC	TGCGCTCCCT	GTACAACACC	1560
ATCGCCGTGC	TGTACTGCGT	GCACCAGCGC	ATCGACGTGA	AGGACACCAA	GGAGGCCCTG	1620
GACAAGATCG	AGGAGGAGCA	GAACAAGTCC	AAGAAGAAGG	CCCAGCAGGC	CGCCGCCGAC	1680
ACCGGCAACA	ACTCCCAGGT	GTCCCAGAAC	TACCCCATCG	TGCAGAACCT	GCAGGGCCAG	1740
	AGGCCATCTC					1800
	CCCCCGAAGT					1860
CAGGACCTGA	ACACCATGCT	GAACACCGTG	GGCGGCCACC	AGGCCGCCAT	GCAGATGCTG	1920
AAGGAGACCA	TCAACGAGGA	GGCCGCCGAG	TGGGACCGCC	TGCACCCCGT	GCACGCCGGC	1980
	CCGGCCAGAT					2040
	AGCAGATCGG					2100
	GGATCATCCT					2160
	TCCGCCAGGG					2220
AAGACCCTGC	GCGCCGAGCA	GGCCTCCCAG	GAGGTAAAGA	ACTGGATGAC	CGAGACCCTG	2280
	ACGCCAACCC					2340
	AGATGATGAC					2400
	AGGCCATGTC					2460
	ACCAGCGCAA					2520
	CCCCCCCCC					2580
	ATTGTACTGA					2640
	CAGGGAATTT					2700
	TTGGGGAAGA					2760
GAACTGTATC	CTTTAGCTTC	CCTCAGATCA	CTCTTTGGCA	GCGACCCCTC	GTCACAATAA	2820

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AGATCGGTGG	CCAGCTGAAG	GAGGCCCTGC	TGGACACCGG	CGCCGACGAC	ACCGTGCTGG	2880
AGGAGATGAA	CCTGCCCGGC	CGCTGGAAGC	CCAAGATGAT	CGGCGGCATC	GGCGGCTTCA	2940
TCAAAGTCCG	CCAGTACGAC	CAGATCCTGA	TCGAGATCTG	CGGCCACAAG	GCCATCGGCA	3000
CCGTGCTGGT	GGGCCCCACC	CCCGTGAACA	TCATCGGCCG	CAACCTGCTG	ACCCAGATCG	3060
GCTGCACCCT	GAACTTCCCC	ATCTCCCCCA	TCGAGACCGT	GCCCGTGAAG	CTGAAGCCCG	3120
GCATGGACGG	CCCCAAAGTC	AAGCAGTGGC	CCCTGACCGA	GGAGAAGATC	AAGGCCCTGG	3180
TGGAGATCTG	CACCGAGATG	GAGAAGGAGG	GCAAGATCTC	CAAGATCGGC	CCCGAGAACC	3240
CCTACAACAC	CCCCGTGTTC	GCCATCAAGA	AGAAGGACTC	CACCAAGTGG	CGCAAGCTGG	3300
TGGACTTCCG	CGAGCTGAAC	AAGCGCACCC	AGGACTTCTG	GGAGGTGCAG	CTGGGCATCC	3360
CCCACCCCGC	CGGCCTGAAG	CAGAAGAAGT	CCGTGACCGT	GCTGGACGTG	GGCGACGCCT	3420
ACTTCTCCGT	GCCCCTGGAC	AAGGACTTCC	GCAAGTACAC	CGCCTTCACC	ATCCCCTCCA	3480
TCAACAACGA	GACCCCCGGC	ATCCGCTACC	AGTACAACGT	GCTGCCCCAG	GGCTGGAAGG	3540
GCTCCCCCGC	CATCTTCCAG	TGCTCCATGA	CCAAGATCCT	GGAGCCCTTC	CGCAAGCAGA	3600
ACCCCGACAT	CGTGATCTAC	CAGTACATGG	ACGACCTGTA	CGTGGGCTCC	GACCTGGAGA	3660
TCGGCCAGCA	CCGCACCAAG	ATCGAGGAGC	TGCGCCAGCA	CCTGCTGCGC	TGGGGCTTCA	3720
CCACCCCGA	CAAGAAGCAC	CAGAAGGAGC	CCCCCTTCCT	GTGGATGGGC	TACGAGCTGC	3780
ACCCCGACAA	GTGGACCGTG	CAGCCCATCG	TGCTGCCCGA	GAAGGACTCC	TGGACCGTGA	3840
ACGACATCCA	GAAGCTGGTG	GGCAAGCTGA	ACTGGGCCTC	CCAGATCTAC	GCCGGCATCA	3900
AAGTCCGCCA	GCTGTGCAAG	CTGCTGCGCG	GCACCAAGGC	CCTGACCGAG	GTGGTGCCCC	3960
TGACCGAGGA	GGCCGAGCTG	GAGCTGGCCG	AGAACCGCGA	GATCCTGAAG	GAGCCCGTGC	4020
ACGGCGTGTA	CTACGACCCC	TCCAAGGACC	TGATCGCCGA	GATCCAGAAG	CAGGGCCAGG	4080
GCCAGTGGAC	CTACCAGATC	TACCAGGAGC	CCTTCAAGAA	CCTGAAGACC	GGCAAATACG	4140
CCCGCATGAA	GGGCGCCCAC	ACCAACGACG	TGAAGCAGCT	GACCGAGGCC	GTGCAGAAGA	4200
TCGCCACCGA	GTCCATCGTG	ATCTGGGGCA	AGACTCCCAA	GTTCAAGCTG	CCCATCCAGA	4260
AGGAGACCTG	GGAGGCCTGG	.TGGACCGAGT	ACTGGCAGGC	CACCTGGATC	CCCGAGTGGG	4320
AGTTCGTGAA	CACCCCCCC	CTGGTGAAGC	TGTGGTACCA	GCTGGAGAAG	GAGCCCATCA	4380
TCGGCGCCGA	GACCTTCTAC	GTGGACGGCG	CCGCCAACCG	CGAGACCAAG	CTGGGCAAGG	4440
CCGGCTACGT	GACCGACCGC	GGCCGCCAGA	AGGTGGTGCC	CCTGACCGAC	ACCACCAACC	4500
AGAAGACCGA	GCTGCAGGCC	ATCCACCTGG	CCCTGCAAGA	CTCCGGCCTG	GAGGTGAACA	4560
TCGTGACCGA	CTCCCAGTAT	GCATTGGGCA	TCATCCAGGC	CCAGCCCGAC	AAGTCCGAGT	4620
CCGAGCTGGT	GTCCCAGATC	ATCGAGCAGC	TGATCAAGAA	GGAGAAGGTG	TACCTGGCCT	4680
GGGTGCCCGC	CCACAAGGGC	ATCGGCGGCA	ACGAGCAGGT	GGACAAGCTG	GTGTCCGCCG	4740
GCATCCGCAA	GGTGCTGTTC	CTGGACGGCA	TCGACAAGGC	CCAGGAGGAG	CACGAGAAGT	4800
ACCACTCCAA	CTGGCGCGCC	ATGGCCTCCG	ACTTCAACCT	GCCCCCGTG	GTGGCCAAGG	4860
AGATCGTGGC	CTCCTGCGAC	AAGTGCCAGC	TGAAGGGCGA	GGCCATGCAC	GGCCAGGTGG	4920
	CGGCATCTGG					4980
TGGCCGTGCA	CGTGGCCTCC	GGCTACATCG	AGGCCGAGGT	GATCCCCGCC	GAGACCGGCC	5040
AGGAGACCGC	CTACTTCCTG	CTGAAGCTGG	CCGGCCGCTG	GCCCGTGAAG	ACCGTGCACA	5100
CCGACAACGG	CTCCAACTTC	ACCTCCACCA	CCGTGAAGGC	CGCCTGCTGG	TGGGCCGGCA	5160
	GTTCGGCATC					5220
	GAAGAAGATC					5280,
	GGCCGTGTTC					5340
	GCGCATCGTG					5400
	CAAGATCCAG					5460
GGAAGGGCCC	CGCCAAGCTG	CTGTGGAAGG	GCGAGGGCGC	CGTGGTGATC	CAGGACAACT	5520
	GGTGGTGCCC					5580
TGGCCGGCGA	CGACTGCGTG	GCCTCCCGCC	AGGACGAGGA	CTAACACATG	GAAAAGATTA	5640

GTAAAACACC	ATAGGCCGCT	CTAGAGGATC	CAAGCTTATC	GATACCGTCG	ACCTCGAGGG	5700
CCCAGATCTA	ATTCACCCCA	CCAGTGCAGG	CTGCCTATCA	GAAAGTGGTG	GCTGGTGTGG	5760
CTAATGCCCT	GGCCCACAAG	TATCACTAAG	CTCGCTTTCT	TGCTGTCCAA	TTTCTATTAA	5820
AGGTTCCTTT	GTTCCCTAAG	TCCAACTACT	AAACTGGGGG	ATATTATGAA	GGGCCTTGAG	5880
CATCTGGATT	CTGCCTAATA	AAAAACATTT	ATTTTCATTG	CAATGATGTA	TTTAAATTAT	5940
TTCTGAATAT	TTTACTAAAA	AGGGAATGTG	GGAGGTCAGT	GCATTTAAAA	CATAAAGAAA	6000
TGAAGAGCTA	GTTCAAACCT	TGGGAAAATA	CACTATATCT	TAAACTCCAT	GAAAGAAGGT	6060 -
GAGGCTGCAA	ACAGCTAATG	CACATTGGCA	ACAGCCCCTG	ATGCCTATGC	CTTATTCATC	6120
CCTCAGAAAA	GGATTCAAGT	AGAGGCTTGA	TTTGGAGGTT	AAAGTTTTGC	TATGCTGTAT	6180
TTTACATTAC	TTATTGTTTT	AGCTGTCCTC	ATGAATGTCT	TTTCACTACC	CATTTGCTTA	6240
TOTTGCATCT	CTCAGCCTTG	ACTCCACTCA	GTTCTCTTGC	TTAGAGATAC	CACCTTTCCC	6300
CTGAACTGTT	CCTTCCATGT	TTTACGGCGA	GATGGTTTCT	CCTCGCCTGG	CCACTCAGCC	6360
TTAGTTGTCT	CTGTTGTCTT	ATAGAGGTCT	ACTTGAAGAA	GGAAAAACAG	GGGGCATGGT	6420
TTCACTCTCC	TGTGAGCCCT	TCTTCCCTGC	CTCCCCCACT	CACAGTGACC	CGGAATCCCT	6480
CCACATGGCA	GTCTAGATCA	TTCTTGAAGA	CGAAAGGGCC	TCGTGATACG	CCTATTTTTA	6540
TACCTTAATC	TCATGATAAT	AATGGTTTCT	TAGACGTCAG	GTGGCACTTT	TCGGGGAAAT	6600
CTCCCCCGGAA	CCCCTATTTG	TTTATTTTTC	TAAATACATT	CAAATATGTA	TCCGCTCATG	6660
AGACAATAAC	CCTGATAAAT	GCTTCAATAA	TATTGAAAAA	GGAAGAGTAT	GAGTATTCAA	6720
CATTTCCGTG	TCGCCCTTAT	TCCCTTTTTT	GCGGCATTTT	GCCTTCCTGT	TTTTGCTCAC	6780
CCAGAAACGC	TGGTGAAAGT	AAAAGATGCT	GAAGATCAGT	TGGGTGCACG	AGTGGGTTAC	6840
ATCGAACTGG	ATCTCAACAG	CGGTAAGATC	CTTGAGAGTT	TTCGCCCCGA	AGAACGTTTT	6900
CCAATGATGA	GCACTTTTAA	AGTTCTGCTA	TGTGGCGCGG	TATTATCCCG	TATTGACGCC	6960
GGGCAAGAGC	AACTCGGTCG	CCGCATACAC	TATTCTCAGA	ATGACTTGGT	TGAGTACTCA	7020
CCAGTCACAG	AAAAGCATCT	TACGGATGGC	ATGACAGTAA	GAGAATTATG	CAGTGCTGCC	7080
ATAACCATGA	GTGATAACAC	TGCGGCCAAC	TTACTTCTGA	CAACGATCGG	AGGACCGAAG	7140
CACCTAACCG	CTTTTTTGCA	CAACATGGGG	GATCATGTAA	CTCGCCTTGA	TCGTTGGGAA	7200
CCCCACCTGA	ATGAAGCCAT	ACCAAACGAC	GAGCGTGACA	CCACGATGCC	TGTAGCAATG	7260
CCAACAACGT	TGCGCAAACT	ATTAACTGGC	GAACTACTTA	CTCTAGCTTC	CCGGCAACAA	7320
TTAATAGACT	GGATGGAGGC	GGATAAAGTT	GCAGGACCAC	TTCTGCGCTC	GGCCCTTCCG	7380
CCTCCCTCCT	TTATTGCTGA	TAAATCTGGA	GCCGGTGAGC	GTGGGTCTCG	CGGTATCATT	7440
GCAGCACTGG	GGCCAGATGG	TAAGCCCTCC	CGTATCGTAG	TTATCTACAC	GACGGGGAGT	7500
CAGGCAACTA	TGGATGAACG	AAATAGACAG	ATCGCTGAGA	TAGGTGCCTC	ACTGATTAAG	7560
CATTGGTAAC	TGTCAGACCA	AGTTTACTCA	TATATACTTT	AGATTGATTT	AAAACTTCAT	7620
TTTTAATTTA	AAAGGATCTA	GGTGAAGATC	CTTTTTGATA	ATCTCATGAC	CAAAATCCCT	7680
TAACGTGAGT	TTTCGTTCCA	CTGAGCGTCA	GACCCCGTAG	AAAAGATCAA	AGGATCTTCT	7740
TGAGATCCTT	TTTTTCTGCG	CGTAATCTGC	TGCTTGCAAA	CAAAAAAAACC	ACCGCTACCA	7800
CCCCTCCTT	GTTTGCCGGA	TCAAGAGCTA	CCAACTCTTT	TTCCGAAGGT	AACTGGCTTC	7860
AGCAGAGCGC	AGATACCAAA	TACTGTTCTT	CTAGTGTAGC	CGTAGTTAGG	CCACCACTTC	7920
AAGAACTCTG	TAGCACCGCC	TACATACCTC	GCTCTGCTAA	TCCTGTTACC	AGTGGCTGCT	7980
CCCACTGGCG	ATAAGTCGTG	TCTTACCGCG	TTGGACTCAA	GACGATAGTT	ACCGGATAAG	8040
GCGCAGCGGT	CGGGCTGAAC	GGGGGGTTCG	TGCACACAGC	CCAGCTTGGA	GCGAACGACC	8100
TACACCGAAC	TGAGATACCT	ACAGCGTGAG	CTATGAGAAA	GCGCCACGCT	TCCCGAAGGG	8160
AGAAAGGCGG	ACAGGTATCC	GGTAAGCGGC	AGGGTCGGAA	CAGGAGAGCG	CACGAGGGAG	8220
CTTCCAGGGG	GAAACGCCTG	GTATCTTTAT	AGTCCTGTCG	GGTTTCGCCA	CCTCTGACTT	8280
GAGCGTCGAT	TTTTGTGATG	CTCGTCAGGG	GGGCGGAGCC	TATGGAAAAA	CGCCAGCAAC	8340
GGATGCGCCG	CGTGCGGCTG	CTGGAGATGG	CGGACGCGAT	GGATATGTTC	TGCCAAGGGT	8400
TGGTTTGCGC	ATTCACAGTT	CTCCGCAAGA	ATTGATTGGC	TCCAATTCTT	GGAGTGGTGA	8460

Fig. 10C

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					TCCATGCACC	. 8520
GCGACGCAAC	GCGGGGAGGC	AGACAAGGTA	TAGGGCGGCG	CCTACAATCC	ATGCCAACCC	8580
GTTCCATGTG	CTCGCCGAGG	CGGCATAAAT	CCCCGTGACG	ATCAGCGGTC	CAATGATCGA	8640
AGTTAGGCTG	GTAAGAGCCG	CGAGCGATCC	TTGAAGCTGT	CCCTGATGGT	CGTCATCTAC	8700
CTGCCTGGAC	AGCATGGCCT	GCAACGCGGG	CATCCCGATG	CCGCCGGAAG	CGAGAAGAAT	8760
CATAATGGGG	AAGGCCATCC	AGCCTCGCGT	CGGGGAGCTT	TTTGCAAAAG	CCTAGGCCTC	8820
	TCCTCACTAC					8880
	AAAAATTACT					

Fig. 10D

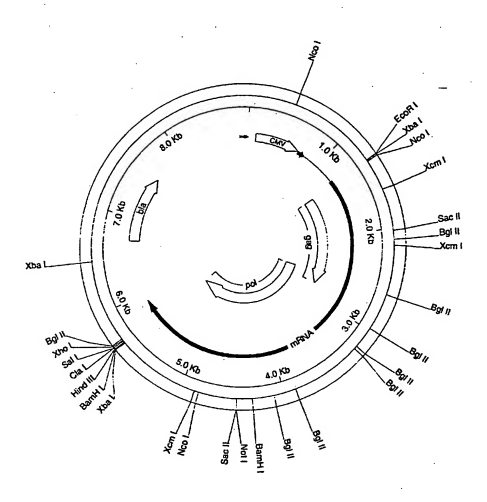


Fig. 11

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A CLASS	FICATION OF SUBJECT C12N15/86	C12N5/10	C12N7/04	C12N15/49	C07K14/16
According to	o International Palent Clas	sification (IPC) or to bo	th national classification	and IPC	
	SEARCHED				
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Documentat	ion searched other than n	ninimum documentation	to the extent that such o	locuments are included in	the fields searched
Electronic d	ata base consulted during	the international searc	th (name of data base an	d, where practical, search	n terms used)
C. DOCUME	ENTS CONSIDERED TO	BE RELEVANT			
Category *	Citation of document, wi	th indication, where ap	propriate, of the relevant	passages	Relevant to claim No.
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			-/		
X Furth	er documents are listed in	the continuation of box	r C.	Patent family member	s are fisted in annex.
"A" documer conside "E" earlier de filling da "L" documen which is citation "O" documen other in "P" documen	it which may throw doubts scited to establish the put or other special reason (a nt referring to an oral disci	te of the art which is no vance or after the internations on priority claim(s) or fleation date of enother a specified) osure, use, exhibition of emational (fling date by	al occupant	or priority date and not in o itseld to understand the pri- mention counterst or particular relav- cannot be considered novi- motive an inventive step w ocument of particular relev- cannot be considered to in- counters of which and the considered of in-	ther the intermetional filing date conflict with the application but notice or theory underlying the vence; the claimed invention of or cannot be concidered to then the document is taken alone ance; the claimed invention volve an inventive stap when the none or more other such docu- eting obvious to a person stolled time patent family
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Name and m	sking address of the ISA European Patent Office	, P.B. 5818 Patentiean		uthorized officer	
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	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	
Category *	Citation of document, with indication where appropriate, of the relevant passages	Relevant to claim No.
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